To: Greaves, Holly[greaves.holly@epa.gov]; Beck, Nancy[Beck.Nancy@epa.gov]; Yamada,

Richard (Yujiro)[yamada.richard@epa.gov]

From: Dourson, Michael

Sent: Thur 11/2/2017 5:27:20 PM

Subject: Re: IRIS/TSCA

Holly, Nancy and Richard

I have talked with a number of senior risk assessment career EPA staff over the years regarding

Ex. 5 - Deliberative Process

I would be more than happy to work up a straw-person for such a group if this is helpful for our discussion tomorrow.

Cheers!

Michael

Sent from my iPad

On Nov 1, 2017, at 10:07 AM, Dourson, Michael < dourson.michael@epa.gov > wrote:

Holly, thanks for the invitation to participate. I am looking forward to it.

Richard, congratulations on getting the boards announced. Very nice.

Cheers!

Michael

From: Greaves, Holly

Sent: Wednesday, November 1, 2017 9:38 AM

To: Dourson, Michael < dourson.michael@epa.gov >; Beck, Nancy

< Beck. Nancy@epa.gov >; Yamada, Richard (Yujiro) < yamada.richard@epa.gov > **Subject:** RE: IRIS/TSCA Good morning, I'd like to re-visit this issue with the 3 of you now that the Boards have been announced. The best time appears to be Friday at 2:30 (Nancy, I see you will have to leave early – if you'd prefer to call-in instead, please let me know). Thanks, Holly From: Dourson, Michael Sent: Tuesday, October 24, 2017 7:31 AM **To:** Greaves, Holly <greaves.holly@epa.gov>; Beck, Nancy <Beck.Nancy@epa.gov>; Yamada, Richard (Yujiro) < yamada.richard@epa.gov> Subject: RE: IRIS/TSCA Holly

Ex. 5 - Deliberative Process

Cheers!

Michael

From: Greaves, Holly

Sent: Monday, October 23, 2017 11:52 AM

To: Dourson, Michael < dourson.michael@epa.gov >; Beck, Nancy

< Beck. Nancy@epa.gov >; Yamada, Richard (Yujiro) < yamada.richard@epa.gov >

Subject: RE: IRIS/TSCA

Dr. Dourson, thank you so much – this is really a helpful starting point.

From the comments below, and my knowledge of the program, the summarized options that we have are as follows:

Ex. 5 - Deliberative Process

Does this group feel that we could take these options to Ryan to obtain his direction on how to move forward? If we do that, it would be helpful to have consensus among this group to provide Ryan with one recommendation from potential options.

Please let me know.

From: Dourson, Michael

Sent: Saturday, October 21, 2017 12:42 PM

To: Beck, Nancy < <u>Beck.Nancy@epa.gov</u>>; Yamada, Richard (Yujiro) < <u>yamada.richard@epa.gov</u>>; Greaves, Holly < <u>greaves.holly@epa.gov</u>>

Subject: Re: IRIS/TSCA

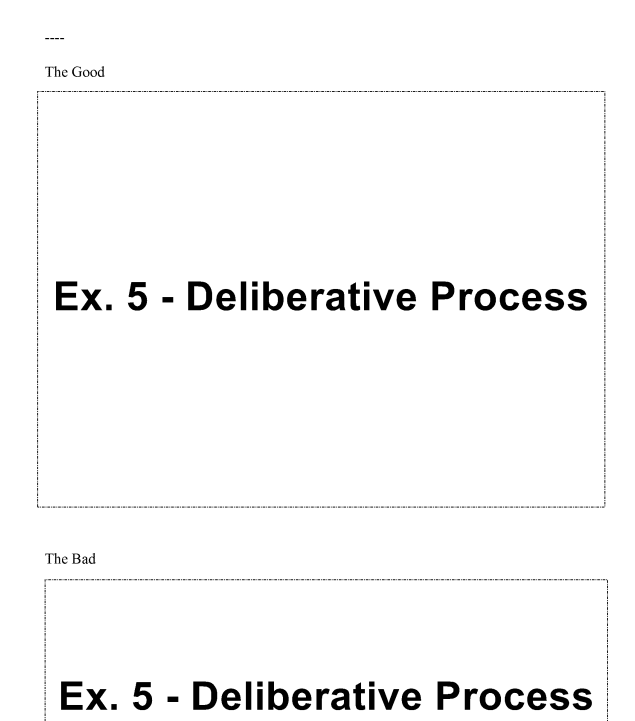
Dear Holly, Nancy and Richard

Thanks for including me in the IRIS discussion. As one of the first IRIS leaders I have been concerned about it ever since the late 1990s. What follows is a sketch that outlines 3 items, roughly described at the good, the bad, and the possible!

I would be more than happy to flesh these musings out along with a lot of input from you and other colleagues in EPA.

Cheers!

Michael



ED 001803B 00000064-5

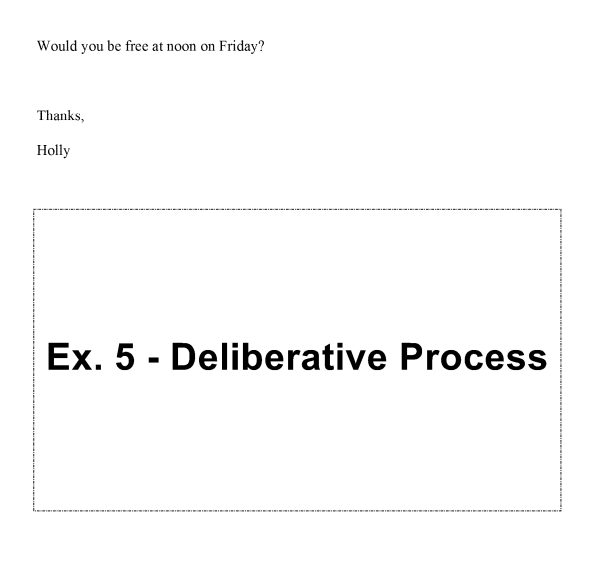
Ex. 5 - Deliberative Process

Sent from my iPad

On Oct 19, 2017, at 7:14 PM, Dourson, Michael < <u>dourson.michael@epa.gov</u> > wrote: Thanks!
From: Beck, Nancy Sent: Thursday, October 19, 2017 7:51 AM To: Dourson, Michael < dourson.michael@epa.gov> Subject: FW: IRIS/TSCA
FYI—background for Friday. I can give you more info as well.
Nancy B. Beck, Ph.D., DABT Deputy Assistant Administrator, OCSPP
P: 202-564-1273
M: Ex. 6 - Personal Privacy beck.nancy@epa.gov
From: Greaves, Holly Sent: Wednesday, October 18, 2017 10:11 AM To: Beck, Nancy < Beck. Nancy@epa.gov >; Yamada, Richard (Yujiro) < yamada.richard@epa.gov > Subject: IRIS/TSCA

Good morning,

The political Associate Director at OMB reached out again today about Ex. 5 - Deliberative Process I'm copying his comments below. I would like to set up a call with him and the 3 of us.



To: Dourson, Michael[dourson.michael@epa.gov]

From: Dourson, Michael

Sent: Sat 11/4/2017 12:35:28 AM

Subject: Fwd: IRIS

Sent from my iPad

Begin forwarded message:

From: dourson.michael@epa.gov

Date: November 3, 2017 at 7:56:30 PM EDT

To: dourson.michael@epa.gov

Subject: IRIS

History

IRIS is a database containing information about a chemical's principle toxic effect and a concentration or dose at which the chemical will not cause any significant effect, even in sensitive humans. For chemicals that cause cancer as the principle effect, this concentration or dose is associated with a very low risk of cancer (usually one chance in a million people). For chemicals that cause some other principle effect (like liver toxicity), this concentration or dose is considered safe. Collectively, these concentrations or doses are referred to as risk values.

The determination of the principle effect is referred to as hazard identification (although other effects at higher concentrations or doses are also described). The determination of these risk values is referred to as dose response assessment.

Up until 1995, IRIS contained risk values on over 500 chemicals and was considered to be the place where all important EPA risk values were placed. Two senior EPA technical groups reviewed all risk values before placing them on IRIS during monthly meetings. Risk values on IRIS were considered to be EPA values and to be used until more appropriate values were developed.

Ex. 5 - Deliberative Process

Political pressures

Ex. 5 - Deliberative Process

One way forward

Ex. 5 - Deliberative Process

Sent from my iPad

To: Dourson, Michael[dourson.michael@epa.gov]

From: Dourson, Michael

Sent: Fri 11/3/2017 11:56:31 PM

Subject: IRIS

History

IRIS is a database containing information about a chemical's principle toxic effect and a concentration or dose at which the chemical will not cause any significant effect, even in sensitive humans. For chemicals that cause cancer as the principle effect, this concentration or dose is associated with a very low risk of cancer (usually one chance in a million people). For chemicals that cause some other principle effect (like liver toxicity), this concentration or dose is considered safe. Collectively, these concentrations or doses are referred to as risk values.

The determination of the principle effect is referred to as hazard identification (although other effects at higher concentrations or doses are also described). The determination of these risk values is referred to as dose response assessment.

Up until 1995, IRIS contained risk values on over 500 chemicals and was considered to be the place where all important EPA risk values were placed. Two senior EPA technical groups reviewed all risk values before placing them on IRIS during monthly meetings. Risk values on IRIS were considered to be EPA values and to be used until more appropriate values were developed.

Ex. 5 - Deliberative Process

Political pressures

Sent from my iPad

Charlotte[Bertrand.Charlotte@epa.gov] From: Dourson, Michael Thur 11/2/2017 1:04:58 PM Sent: Subject: Endangered Species Act Options Rick, Nancy and Charlotte The scientists in OPP on this issue are at the top of their game. Unfortunately, we did not have a face-to-face meeting with our DOI colleagues yesterday, but both Kris and Melissa had enough time to gain a sense on what the Biological Opinions were based. After walking back with them in discussion, I got a better sense of our options. I list these below. Your thoughts on these would be extremely valuable. Cheers! Michael Options for ESA work Ex. 5 - Deliberative Process

Keigwin, Richard[Keigwin.Richard@epa.gov]; Beck, Nancy[beck.nancy@epa.gov]; Bertrand,

To:

Dourson, Michael[dourson.michael@epa.gov] Michael Dourson To:

From:

Sent: Mon 1/22/2018 2:16:18 PM

Subject: Testing

Cc: Beck, Nancy[Beck.Nancy@epa.gov]; Hanley, Mary[Hanley.Mary@epa.gov]; Wise, Louise[Wise.Louise@epa.gov]; Bertrand, Charlotte[Bertrand.Charlotte@epa.gov]; Keller, Kaitlin[keller.kaitlin@epa.gov]; Jakob, Avivah[Jakob.Avivah@epa.gov]; Bolen, Derrick[bolen.derrick@epa.gov] To: Kaiser, Sven-Erik[Kaiser.Sven-Erik@epa.gov] From: Dourson, Michael Sent: Tue 10/24/2017 9:28:24 PM Subject: Re: SEPW Minority Letter to Dr. Dourson Dear Mr. Kaiser I would be happy to assist in this response. Several of these questions can be easily handled. Please let me know of when you need a draft. Cheers! Michael Dourson Sent from my iPad > On Oct 24, 2017, at 4:52 PM, Kaiser, Sven-Erik <Kaiser, Sven-Erik@epa.gov> wrote: > OCSPP Team - thanks for handling. For reference, here's a similar exchange regarding Susan Bodine (incoming, response and attachments included). Please let me know if any questions. Best, > Sven > Sven-Erik Kaiser > U.S. EPA > Office of Congressional and Intergovernmental Relations > 1200 Pennsylvania Ave., NW (1305A) > Washington, DC 20460 > 202-566-2753 > From: Beck, Nancy > Sent: Tuesday, October 24, 2017 4:45 PM > To: Hanley, Mary <Hanley.Mary@epa.gov>; Wise, Louise <Wise.Louise@epa.gov>; Bertrand, Charlotte <Bertrand.Charlotte@epa.gov>; Keller, Kaitlin <keller.kaitlin@epa.gov>; Jakob, Avivah <Jakob.Avivah@epa.gov> > Cc: Dourson, Michael <dourson.michael@epa.gov>; Bolen, Derrick <bolen.derrick@epa.gov>; Kaiser, Sven-Erik <Kaiser.Sven-Erik@epa.gov> > Subject: FW: SEPW Minority Letter to Dr. Dourson > Mary, > Can you take the lead on getting a response drafted? We will likely need assistance from OCIR, OGC and OPPT. > Draft by next Friday? Is that possible? > Thanks. > Nancy B. Beck, Ph.D., DABT > Deputy Assistant Administrator, OCSPP > P: 202-564-1273 > Mi Ex. 6 - Personal Privacy

> beck.nancy@epa.gov<mailto:beck.nancy@epa.gov>

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> From: Kaiser, Sven-Erik
> Sent: Tuesday, October 24, 2017 3:54 PM
> To: Hanley, Mary <Hanley.Mary@epa.gov<mailto:Hanley.Mary@epa.gov>>; Beck, Nancy
<Beck.Nancy@epa.gov<mailto:Beck.Nancy@epa.gov>>; Wise, Louise
<Wise.Louise@epa.gov<mailto:Wise.Louise@epa.gov>>; Bertrand, Charlotte
<Bertrand.Charlotte@epa.gov<mailto:Bertrand.Charlotte@epa.gov>>; Jakob, Avivah
<Jakob.Avivah@epa.gov<mailto:Jakob.Avivah@epa.gov>>; Keller, Kaitlin
<keller.kaitlin@epa.gov<mailto:keller.kaitlin@epa.gov>>
> Subject: SEPW Minority Letter to Dr. Dourson
> OCSPP Team - heads up on a letter to Dr. Dourson. I'm checking with OCIR management on handling
and will let you know as soon as I hear something. Please let me know if any questions. Thanks,
> Sven
> Sven-Erik Kaiser
> U.S. EPA
> Office of Congressional and Intergovernmental Relations
> 1200 Pennsylvania Ave., NW (1305A)
> Washington, DC 20460
> 202-566-2753
> From: Lyons, Troy
> Sent: Tuesday, October 24, 2017 1:14 PM
> To: Aarons, Kyle <Aarons.Kyle@epa.gov<mailto:Aarons.Kyle@epa.gov>>; Palich, Christian
<palich.christian@epa.gov<mailto.palich.christian@epa.gov>>, Kaiser, Sven-Erik <Kaiser.Sven-</p>
Erik@epa.gov<mailto:Kaiser.Sven-Erik@epa.gov>>
> Subject: FW: Letter to Dr. Michael Dourson
> From: Ferrato, Margaret (Whitehouse) [mailto:Margaret Ferrato@whitehouse.senate.gov]
> Sent: Tuesday, October 24, 2017 1:04 PM
> To: Lyons, Troy <lyons.troy@epa.gov<mailto:lyons.troy@epa.gov>>
> Cc: Gaeta, Joe (Whitehouse)
<Joe Gaeta@whitehouse.senate.gov<mailto:Joe Gaeta@whitehouse.senate.gov>>; Leibman, Adena
(Whitehouse)
<Adena Leibman@whitehouse.senate.gov<mailto:Adena Leibman@whitehouse.senate.gov>>; Goldner,
Aaron (Whitehouse)
<Aaron Goldner@whitehouse.senate.gov<mailto:Aaron Goldner@whitehouse.senate.gov>>
> Subject: Letter to Dr. Michael Dourson
> Hi Troy,
> I hope you're well! Attached is a letter from members of the Environment and Public Works Committee
to Dr. Dourson. Don't hesitate to reach out with any questions.
> Best,
> Maggie
> <position description.pdf>
> <signed pledge.pdf>
> <Whitehouse Itr 9-22-17.pdf>
> <Whitehouse Merkley 9-21-17 (Bodine).docx>
> <Whitehouse Merkley Bodine 9-13-17.pdf>
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To: Garber, Kristina[Garber.Kristina@epa.gov]; Panger, Melissa[Panger.Melissa@epa.gov] Anderson, Brian[Anderson.Brian@epa.gov]; Bolen, Derrick[bolen.derrick@epa.gov]; Keigwin, Cc: Richard[Keigwin.Richard@epa.gov]; Keller, Kaitlin[keller.kaitlin@epa.gov]; Beck, Nancy[beck.nancy@epa.gov]; Bertrand, Charlotte[Bertrand.Charlotte@epa.gov] From: Dourson, Michael Thur 11/2/2017 12:45:51 PM Sent: Subject: Endangered Species Dear Kris and Melissa Thanks so much for going over to DOI yesterday. Although we did not get the FTF discussion that we (or at least I) expected, the visit was nevertheless fruitful, due entirely to your collective knowledge of the issues. I will writing a polite note back to David Barnhardt later this morning and making arrangements to visit NOA. Hopefully you can both attend this meeting as well, although it may have to be via Skype.

Sometime very soon, we will have to brief the OCSPP upper management.

Cheers!

Michael...

... L. Dourson, PhD., DABT, FATS, FSRA

Senior Advisor to the Administrator

U.S. Environmental Protection Agency

dourson.michael@epa.gov

202-564-2463

www.epa.gov

----Original Appointment----

From: Bolen, Derrick

Sent: Friday, October 27, 2017 5:44 PM

To: Bolen, Derrick; Dourson, Michael; Garber, Kristina; Panger, Melissa; Anderson, Brian

Subject: HOLD: DOI

When: Wednesday, November 1, 2017 3:00 PM-5:00 PM (UTC-05:00) Eastern Time (US &

Canada). Where: DOI

To: Fugh, Justina[Fugh.Justina@epa.gov]

From: Dourson, Michael

Sent: Tue 1/2/2018 2:49:23 PM

Subject: RE: CAUTION: You're getting closer to being added to the ethics naughty list

Thanks!

From: Fugh, Justina

Sent: Tuesday, January 2, 2018 9:37 AM

To: Dourson, Michael <dourson.michael@epa.gov>

Subject: RE: CAUTION: You're getting closer to being added to the ethics naughty list

Hi there,

Please disregard this notification. We know you already had in-person training. We just neglected to update our database with the date of your training. Sorry about that.

Justina

Justina Fugh | Senior Counsel for Ethics | Office of General Counsel | US EPA | Mail Code 2311A | Room 4308 North, William Jefferson Clinton Federal Building | Washington, DC 20460 (for ground deliveries, use 20004 for the zip code) | phone 202-564-1786 | fax 202-564-1772

From: Dourson, Michael

Sent: Saturday, December 30, 2017 10:21 AM **To:** Fugh, Justina < Fugh. Justina@epa.gov >

Subject: Re: CAUTION: You're getting closer to being added to the ethics naughty list

Justina

I will be working the week of Jan 2, but this may be my last week. This, I hope you do not mind if I forgo the training. If I stay on, I will take the training. If not, then I will need to talk with you about future work.

Holiday cheers!

Michael

Sent from my iPhone

On Dec 19, 2017, at 11:06 AM, Fugh, Justina < Fugh. Justina@epa.gov > wrote:

A gentle nudge – please take your 2017 annual ethics training before 12/31/17

(and, if you read through this whole message, there's a treat at the end) ...

This year's course -- "Follow the Money – Gifts and Travel" -- examines ethics considerations related to ... you guessed it: gifts and travel! This course (the last one created by Dan Fort) will give employees an overview of the federal ethics requirements and then dive into the do's and don'ts of giving and receiving gifts, including the gift of travel. If you have questions about the course, then please contact our new training officer, Margaret Ross (whom we are delighted to have join us).

Who needs to take this training?

YOU have to take the training because you file the public financial disclosure report and, according to our records, have not yet completed training.

What is the deadline for completing the training?

Training must be completed by December 31, 2017. We know, we're tardy in reminding you, but we have faith in you!

How do you access the training?

To access the training, click on this link: <u>2017 Annual Ethics Training</u> or cut and paste the following address into your browser:

http://intranet.epa.gov/ogc/2017ethicstraining/10.html. This training is still not yet hosted in Skillport; rather, it's on the OGC/ethics intranet site. If you don't have access to the intranet (sigh), then please contact Margaret Ross at ross.margaret@epa.gov.

How do you document completion of the course?

At the end of the training, you will see a screen that directs you to enter your EPA email address (i.e., lastname@epa.gov) in order to certify completion of the course. What will happen is that the system will generate a certificate in your browser, then send you an email confirming your completion, and will add your name to the tracking database in Lotus Notes. Yes, we know that Lotus Notes is going away, but OGC/Ethics will keep our subscription going into next year (and encourage ethics officials to do the same). And before our friends in OEI get upset, yes, we know that we need to move to another system next year.

What if I don't get a training certificate?

You know, if you tell me that you took the training but didn't get a certificate, well, I'll believe you. I mean, would you really lie to me about having taken *ethics* training? Just send me an email with the date you took the training, and I'll generate a certificate for you.

Why isn't the training available on Skillport?

Well, it just isn't...but it will be next year. Until then, we're going to muddle through with the Lotus Notes training tracker this last time.

How can I send feedback and comments?

As you know, our training courses are created entirely by the OGC ethics team, principally through the creative genius of Dan Fort, who recently retired. If you have any comments about the content of this course, or want to provide input about future courses, please contact Margaret Ross (our new ethics training officer) at ross.margaret@epa.gov.

PS – and now the "treat" as promised. We hear that the National Executive Leadership Development Conference (NELDC, colloquially known as the SES Forum) will be held in late January 2018. Provided you complete your online training in 2017, then you will be able to get credit for you 2018 training by attending the conference! You know that our ethics training is fun when it's in person, so please plan to stay for that!

Thanks for your attention to ethics issues! Call us anytime (or email us at ethics@epa.gov).

Justina

Justina Fugh | Senior Counsel for Ethics | Office of General Counsel | US EPA | Mail Code 2311A | Room 4308 North, William Jefferson Clinton Federal Building | Washington, DC 20460 (for ground deliveries, use 20004 for the zip code) | phone 202-564-1786 | fax 202-564-1772

Cc: Bolen, Derrick[bolen.derrick@epa.gov]; Beck, Nancy[Beck.Nancy@epa.gov]

To: Washington, Valerie[Washington.Valerie@epa.gov]

From: Dourson, Michael

Sent: Tue 10/31/2017 8:56:50 PM

Subject: Car for transport to Department of Interior

Valerie

Would you please be so kind and work with Derrick Bowen on getting a car for me to take to the Department of Interior tomorrow? Derrick has details of the arrangements.

Thanks!

Michael

Sent from my iPad

To: Keigwin, Richard[Keigwin.Richard@epa.gov]

From: Dourson, Michael

Sent: Thur 11/2/2017 12:28:01 PM Subject: RE: NYT Article on Dicamba

Rick

Thanks.

Michael

From: Keigwin, Richard

Sent: Thursday, November 2, 2017 4:57 AM

To: Bertrand, Charlotte <Bertrand.Charlotte@epa.gov>; Beck, Nancy <Beck.Nancy@epa.gov>; Dourson, Michael <dourson.michael@epa.gov>; Strauss, Linda <Strauss.Linda@epa.gov>; Dinkins, Darlene <Dinkins.Darlene@epa.gov>; Goodis, Michael <Goodis.Michael@epa.gov>;

Baris, Reuben <Baris.Reuben@epa.gov>

Subject: NYT Article on Dicamba

https://mobile.nytimes.com/2017/11/01/business/soybeans-pesticide.html?referer=https://www.google.com/

Rick Keigwin

Director, Office of Pesticide Programs

U.S. Environmental Protection Agency

Phone: 703-305-7090

Website: www.epa.gov/pesticides

Sent from my iPhone

To: Beck, Nancy[Beck.Nancy@epa.gov]

Cc: Bertrand, Charlotte[Bertrand.Charlotte@epa.gov]

From: Dourson, Michael

Sent: Thur 12/14/2017 9:47:57 PM

Subject: Re: meet earlier

Stop by the party. It's important. I have to leave at 5:30

Sent from my iPad

On Dec 14, 2017, at 4:01 PM, Beck, Nancy < Beck. Nancy @epa.gov > wrote:

I may have a meeting with Ryan but it isn't confirmed yet. If I don't here from him can we we do 4:45? I'd like to pop by the OPPT party.

Just leaving HHS now.

Thanks.

Nancy B. Beck, Ph.D., DABT Deputy Assistant Administrator, OCSPP

P: 202-564-1273
M: Ex. 6 - Personal Privacy
Beck. Nancy@epa.gov

On Dec 14, 2017, at 2:48 PM, Bertrand, Charlotte < Bertrand. Charlotte@epa.gov > wrote:

4:30 works for me – Nancy?

From: Dourson, Michael

Sent: Thursday, December 14, 2017 2:48 PM

To: Beck, Nancy < Beck. Nancy@epa.gov >; Bertrand, Charlotte

<Bertrand.Charlotte@epa.gov>

Subject: meet earlier

Nancy and Charlotte

Can we meet earlier than 5 pm today? I have a 6 pm dinner at a metro stop a wee bit away.

Cheers!

Michael...

... L. Dourson, PhD., DABT, FATS, FSRA
Senior Advisor to the Administrator

U.S. Environmental Protection Agency

dourson.michael@epa.gov

202-564-2463

www.epa.gov

To: Beck, Nancy[beck.nancy@epa.gov]; Bertrand, Charlotte[Bertrand.Charlotte@epa.gov]

From: Dourson, Michael

Sent: Wed 11/22/2017 2:07:03 PM

Subject: RE: Chlorpyrifos epidemiology studies

Yes, you have my schedule and I have some thoughts to share.

From: Beck, Nancy

Sent: Tuesday, November 21, 2017 7:39 PM

To: Dourson, Michael <dourson.michael@epa.gov>; Bertrand, Charlotte

<Bertrand.Charlotte@epa.gov>

Subject: RE: Chlorpyrifos epidemiology studies

Chat tomorrow?

Nancy B. Beck, Ph.D., DABT

Deputy Assistant Administrator, OCSPP

P: 202-564-1273

M: Ex. 6 - Personal Privacy

beck.nancy@epa.gov

From: Dourson, Michael

Sent: Tuesday, November 21, 2017 11:29 AM

To: Beck, Nancy <Beck.Nancy@epa.gov>; Bertrand, Charlotte <Bertrand.Charlotte@epa.gov>

Subject: RE: Chlorpyrifos epidemiology studies

Nancy and Charlotte

I have some thoughts on this. Please call to clarify when appropriate.

Cheers!
Michael
L. Dourson, PhD., DABT, FATS, FSRA
Senior Advisor to the Administrator
U.S. Environmental Protection Agency
dourson.michael@epa.gov
202-564-2463
www.epa.gov
Ex. 6 - Personal Privacy
From: Beck, Nancy Sent: Monday, November 20, 2017 6:25 PM To: Dourson, Michael dourson.michael@epa.gov >; Bertrand, Charlotte Bertrand.Charlotte@epa.gov > Subject: RE: Chlorpyrifos epidemiology studies
Ok. How many articles is that? Is there one key one or a few key ones she should focus on? The more specific we can be, the better?
Thanks.

Nancy B. Beck, Ph.D., DABT

Deputy Assistant Administrator, OCSPP

P: 202-564-1273

M: Ex. 6 - Personal Privacy

beck.nancy@epa.gov

From: Dourson, Michael

Sent: Monday, November 20, 2017 3:28 PM

To: Beck, Nancy < Beck. Nancy@epa.gov >; Bertrand, Charlotte < Bertrand. Charlotte@epa.gov >

Subject: RE: Chlorpyrifos epidemiology studies

Nancy

Ex. 5 - Deliberative Process

Cheers!

Michael

From: Beck, Nancy

Sent: Monday, November 20, 2017 2:54 PM

To: Dourson, Michael <<u>dourson.michael@epa.gov</u>>; Bertrand, Charlotte

<Bertrand.Charlotte@epa.gov>

Subject: FW: Chlorpyrifos epidemiology studies

Apologies for coming late to the party, but have we asked OPP to engage OPPT epidemiologists in some way? One person is being bombarded and I think it would help if we could narrow scope/purpose.

See below and lets chat.

Nancy B. Beck, Ph.D., DABT

Deputy Assistant Administrator, OCSPP

P: 202-564-1273

M: Ex. 6 - Personal Privacy

beck.nancy@epa.gov

From: Morris, Jeff

Sent: Monday, November 20, 2017 2:52 PM
To: Beck, Nancy < Beck. Nancy@epa.gov >
Subject: Fwd: Chlorpyrifos epidemiology studies

What's our objective here? Thanks.

Sent from my iPhone

Begin forwarded message:

From: "Henry, Tala" < Henry. Tala@epa.gov > Date: November 20, 2017 at 1:04:56 PM EST

To: "Morris, Jeff" < Morris.Jeff@epa.gov >, "Mottley, Tanya" < Mottley.Tanya@epa.gov >

Subject: FW: Chlorpyrifos epidemiology studies

Jeff/Tanya,

Ex. 5 - Deliberative Process

Please advise.

Tala R. Henry, Ph.D.

Director, Risk Assessment Division

Office of Pollution Prevention and Toxics

U.S. Environmental Protection Agency

T: 202-564-2959

E: henry.tala@epa.gov

From: Pfahles-Hutchens, Andrea

Sent: Monday, November 20, 2017 12:31 PM **To:** Henry, Tala Henry.Tala@epa.gov>

Cc: Laessig, Susan < <u>Laessig.Susan@epa.gov</u>> Subject: Fw: Chlorpyrifos epidemiology studies

Importance: High

Tala,

I have 5 emails full of studies. Can you please tell me what is going on?

Andrea
Andrea Pfahles-Hutchens
Epidemiologist
US EPA
(202)564-7601
From: Hughes, Hayley Sent: Monday, November 20, 2017 12:12 PM To: Keller, Kaitlin; Pfahles-Hutchens, Andrea Subject: RE: Chlorpyrifos epidemiology studies
Hello Kaitlin,
Thank you for your email. For clarification, are we expected to be prepared to discuss the findings of the studies by next week or is this a preliminary meeting?
Thanks,
Hayley
From: Keller, Kaitlin Sent: Monday, November 20, 2017 11:48 AM To: Hughes, Hayley < hughes.hayley@epa.gov >; Pfahles-Hutchens, Andrea < Pfahles-Hutchens.Andrea@epa.gov >

Subject: Chlorpyrifos epidemiology studies	
Hi Hayley and Andrea,	
As I think you are aware, Nancy and Charlotte would like a meeting with you to discuss the Chlorpyrifos epi studies, Ex. 5 - Deliberative Process Ex. 5 - Deliberative Process	he
First, attached is the 2016 OPP systematic review of all of the studies on neurodevelopmental effects of organophosphate pesticides, as well as the last three Chlorpyrifos SAP reports.	
The following four additional emails will include: Ex. 5 - Deliberative Process Ex. 5 - Deliberative Process	
I'm working to get this on the calendar for early next week, stay tuned for the invite. For OPPT, I'll cc Tala Henry and Jeff Morris on the invite, and for OPP, I'll cc Rick Keigwin. Please let me know if anyone else should be aware, or if you have any questions.	
Thanks,	
Kaitlin	
Kaitlin Keller, Special Assistant	
Office of Chemical Safety and Pollution Prevention	
U.S. Environmental Protection Agency	
(202) 564-7098	

To: Washington, Valerie[Washington.Valerie@epa.gov]; Wooden-Aguilar, Helena[Wooden-

Aguilar.Helena@epa.gov]; Allen, Reginald[Allen.Reginald@epa.gov]; Greenwalt,

Sarah[greenwalt.sarah@epa.gov]

From: Dourson, Michael

Sent: Wed 11/15/2017 11:38:47 AM

Subject: RE: Out today

Valerie

Ex. 6 - Personal Privacy

Michael

----Original Message-----From: Washington, Valerie

Sent: Wednesday, November 15, 2017 6:13 AM

To: Wooden-Aguilar, Helena < Wooden-Aguilar. Helena@epa.gov>; Allen, Reginald

<a href="mailto: , Greenwalt, Sarah <a href="mailto: , Dourson, Michael

<dourson.michael@epa.gov>

Subject: Out today

Gm All,

Sent from my iPhone

To: Fugh, Justina[Fugh.Justina@epa.gov]

From: Dourson, Michael

Sent: Sat 12/2/2017 12:58:34 AM

Subject: FW: SRA Annual Meeting Plenary Session Monday morning

Justina

So I will be going to this annual Society for Risk Analysis meeting and likely invited to hospitalities where all folks are offered food. What is your call on this please?

Cheers!

Michael...

... L. Dourson, PhD., DABT, FATS, FSRA

Senior Advisor to the Administrator

U.S. Environmental Protection Agency

dourson.michael@epa.gov

202-564-2463

www.epa.gov

From: Dourson, Michael

Sent: Tuesday, November 28, 2017 7:30 PM **To:** Jackson, Ryan jackson.ryan@epa.gov

Cc: Lyons, Troy
Lyons.troy@epa.gov>; Fugh, Justina <Fugh.Justina@epa.gov>; Bertrand,

Charlotte <bertrand.charlotte@epa.gov>; Beck, Nancy <beck.nancy@epa.gov> Subject: FW: SRA Annual Meeting Plenary Session Monday morning</beck.nancy@epa.gov></bertrand.charlotte@epa.gov>
Ryan
I have a pending talk at the upcoming Society for Risk Analysis meeting in Crystal City, on December 11. The topic of the talk is shown in the emails below, but basically is me giving a few slides (5 at most) on risk analysis as an obsolete profession (or not). I am definitely in the "or not" camp. This commitment was made over 6 months ago.
At this point I am listed on the program as my EPA title below. Please advise if you need for me to change anything.
Cheers!
Michael
L. Dourson, PhD., DABT, FATS, FSRA
Senior Advisor to the Administrator
U.S. Environmental Protection Agency
dourson.michael@epa.gov
202-564-2463
www.epa.gov

Sent: Monday, November 27, 2017 11:39 AM To: 'Terje Aven' < terje.aven@uis.no >; Dourson, Michael < dourson.michael@epa.gov > Subject: RE: SRA Annual Meeting Plenary Session Monday morning Agreed, I know Dr. Dourson has an excellent presentation related to risk analysis (or risk assessment) certification, so some discussion of this would be great. From: Terje Aven [mailto:terje.aven@uis.no] Sent: Monday, November 27, 2017 8:52 AM To: Dourson, Michael Cc: Pamela Williams Subject: SV: SRA Annual Meeting Plenary Session Monday morning Thanks a lot Michael, this is excellent, perhaps you can also think about what we should then do to meet this challenge. I know you would highlight training .. Best Terje Sendt fra E-post for Windows 10 Fra: Dourson, Michael < dourson.michael@epa.gov> Sendt: Monday, November 27, 2017 3:56:12 PM Til: Terje Aven **Kopi:** Pamela Williams Emne: RE: SRA Annual Meeting Plenary Session Monday morning Terje

From: Pamela Williams [mailto:pwilliams@erisksciences.com]

Thanks for the gentle reminder. I am of the mind to discuss the misunderstanding of our profession by unskilled folks, and the plethora of opinions, masquerading as erudite, flooding the market, so to speak. We are not obsolete, as much as we are emulated, unfortunately by folks who really do not understand the underlying science.

I will likely have a few slides as examples. I am thinking of a periodic table chart of chemical contaminants in various folks' bodies, and/or the blogs on various synthetic pesticides on our food, meanwhile ignoring, or more likely being ignorant of, the overwhelming proportion of pesticides in food that are naturally occurring.

I very much appreciate your efforts to pull this together and the initial slides from both you and Pamela.

Cheers!?

Michael

From: Terje Aven [mailto:terje.aven@uis.no]
Sent: Friday, November 24, 2017 4:58 AM

To: Dourson, Michael < dourson.michael@epa.gov > Cc: Pamela Williams < pwilliams@erisksciences.com >

Subject: VS: SRA Annual Meeting Plenary Session Monday morning

Hi Michael.

How are things going concerning the preparation for the Panel?

I know we are a little early, but we very much would appreciate some feedback before the end of the month to be able to plan the discussion in a good way.

Thanks a lot

Best

Terje

Fra: Terje Aven

Sendt: 20. oktober 2017 11:31

Til: doursoml@ucmail.uc.edu; ragnar.lofstedt@kcl.ac.uk; sguikema@umich.edu;

kimt@aorm.com

Kopi: Pamela Williams < <u>pwilliams@erisksciences.com</u>>

Emne: SRA Annual Meeting Plenary Session Monday morning

Hi all,

Thanks for participating in the panel **Risk Analysis: An Obsolete Profession?** It will be great :-)

I will have an introduction to the panel discussion, see enclosed preliminary slides with associated text (the last slides 16-25 are not planned to be presented).

After this introduction I give the word to Pamela, see her preliminary slides (not all of these will be used but they are included to make the presentation understandable).

The idea is that each of you has a prepared introduction of some 5-7 minutes, prepared with slides if you like, with clear statements —linked to abstract of the panel and hopefully inspired by mine and Pamela's slides.

We would not like to restrict creativity and what you find most important on this matter, so feel free to angle things in your way. Focusing on some few – one or two – themes is however recommended. To be able to lead the panel discussion in a good way, we think it is wise to have a process in advance – starting now – where we share some of the ideas we have. The aim of this dialogue is to make the panel as interesting as possible by being informed what is coming, so that one can get ideas for comments and questions. We would like to have a lively discussion so the point is not use this dialogue to obtain some unity or consensus at this stage (rather the opposite ③)

Looking forward to hearing from you. What we ask from you now is an indication of what type of message – themes- that you would like to highlight - in text or using slides. We would very much appreciate if we could get some input before 15 November. Thanks a lot. Enjoy the weekend. Best Terje SRA Annual Meeting Plenary sessions Monday morning Risk Analysis: An Obsolete Profession?

Risk analysis has advanced strongly the last 30-40 years. It is interdisciplinary in its scope but also developing as a science in itself. Yet we should ask, has it really evolved as it should? Is there a potential for reaching another level on both quality

and outreach?

Is there a need for revitalization and new directions for the field and SRA, to strengthen the research and reflect current topics like resilience and security? Should we develop specific risk analysis certificates and educational programs?

The panel will discuss these topics - the role of risk analysis in society and how risk analysis as a field can be strengthened. We question, what does it really mean to be a risk analysis practitioner, professional and scientist?

Panel:

Chairs: Terje Aven and Pamela Williams

Michael Dourson, Seth Guikema, Ragnar Löfstedt, Kimberly Thompson

Terje Aven, University of Stavanger, Norway

Pamela Williams, E Risk Sciences

Michael Dourson, US Environmental Protection Agency (EPA) (waiting for final confirmation)

Seth Guikema, University of Michigan

Ragnar Löfstedt, Kings College, London

Kimberly Thompson, Kid Risk and University of Central Florida

Ex. 6 - Personal Privacy To:

From: ⁱ⊔ourson, iviichaei

Sent: Mon 1/1/2018 9:25:02 PM Subject: Fwd: Check in for your flight

Sent from my iPhone

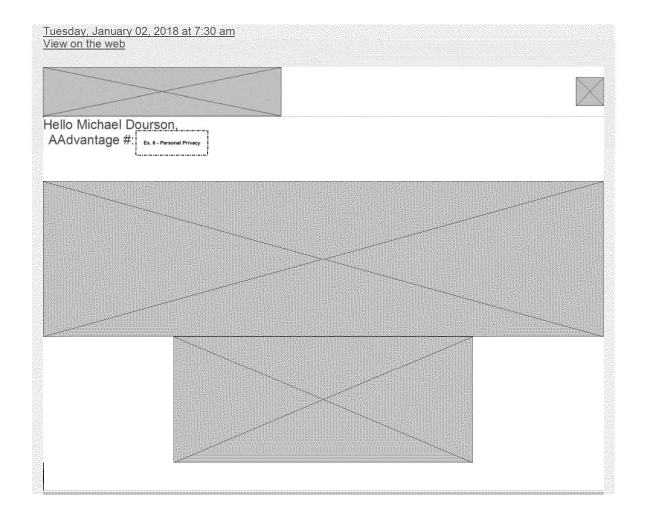
Begin forwarded message:

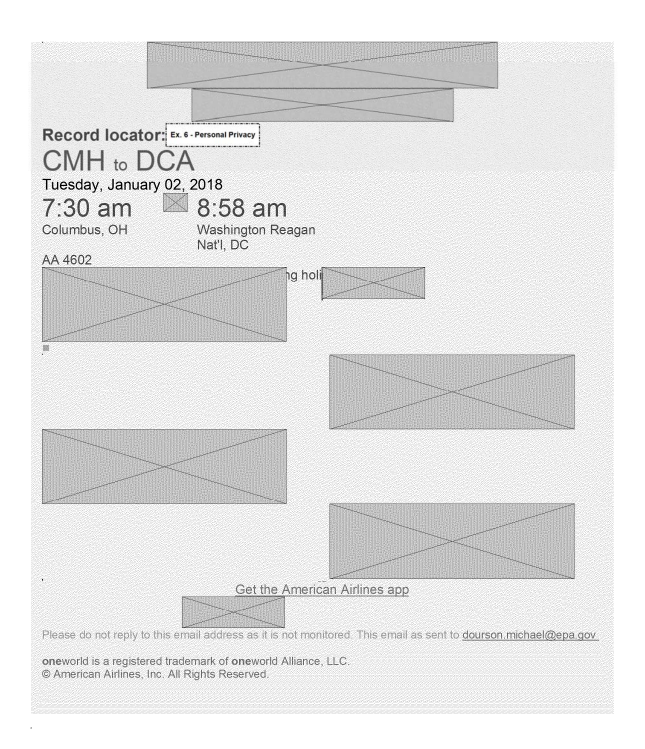
From: "American Airlines" < americanairlines@checkin.email.aa.com>

Date: January 1, 2018 at 12:02:43 PM EST

To: <dourson.michael@epa.gov> Subject: Check in for your flight **Reply-To:** American Airlines

<a href="mailto:mailto:<a href="mailto:255.6701837@checkin.emailto:255.6701837@checkin.emailto:255.6701837@checkin.emailto:255.6701837@checkin.emailto:255.6701837@checkin.e





Subject: Re: National TRI Analysis Release Nancy Very nice email! Mike Sent from my iPad > On Nov 30, 2017, at 12:07 PM, Beck, Nancy <Beck.Nancy@epa.gov> wrote: > I have an opportunity for you all! > Our TRI team (Toxic Release Inventory) (now located in OCSPP, used to be in OEI), is getting ready to the 2016 National Analysis Release in mid-January. The TRI release, while predominantly a good news story, always seems to raise some issues of concerns with certain stakeholder sectors. The way the information is rolled out on the web, there are many tools that can be used for finding all sorts of release information for specific sectors. For instance, while air releases have decreased 58% since 2006, total production of waste increased 2% in 2015-2016 and total land disposal increased 1% (mostly due to metal mining), and lead releases increased 16% (also due to metal mining). These are just some snippets from the attached powerpoint. While land disposal is often to enclosed structures, with no releases to the environment, communicating this information is sometimes challenging. > Liz- I presume OPA may want a briefing? and perhaps the Administrators office as well. Since there are releases identified to air, water, land, etc other programs may have some equities here. > Feel free to a look at the attached powerpoint to get a sense of what the release will look like and please let Derrick Bolen know if you are interested in a briefing. We can work to set something up before the release in January. > https://www.epa.gov/toxics-release-inventory-tri-program > Regards, > Nancv > Nancy B. Beck, Ph.D., DABT > Deputy Assistant Administrator, OCSPP > P: 202-564-1273 > M: Ex. 6 - Personal Privacy > beck.nancy@epa.gov<mailto:beck.nancy@epa.gov> > From: Ford, Hayley > Sent: Wednesday, November 29, 2017 6:44 PM > To: Ford, Hayley <ford.hayley@epa.gov>; McMurray, Forrest <mcmurray.forrest@epa.gov>; Daniell, Kelsi <daniell.kelsi@epa.gov>; Bowman, Liz <Bowman.Liz@epa.gov>; Falvo, Nicholas <falvo.nicholas@epa.gov>; Dravis, Samantha <dravis.samantha@epa.gov>; Greaves, Holly <greaves.holly@epa.gov>; Gunasekara, Mandy <Gunasekara.Mandy@epa.gov>; Chmielewski, Kevin <chmielewski.kevin@epa.gov>; Munoz, Charles <munoz.charles@epa.gov>; Hewitt, James <hewitt.james@epa.gov>; Ringel, Aaron <ringel.aaron@epa.gov>; Frye, Tony (Robert) <frye.robert@epa.gov>; Burke, Marcella <burke.marcella@epa.gov>; Cory, Preston (Katherine) <Cory.Preston@epa.gov>; Palich, Christian <palich.christian@epa.gov>; Ferguson, Lincoln <ferguson.lincoln@epa.gov>; Wilcox, Jahan <wilcox.jahan@epa.gov>; Feeley, Drew (Robert) <Feeley.Drew@epa.gov>; Gordon, Stephen <gordon.stephen@epa.gov>; Dourson, Michael

To:

From:

Sent:

Beck, Nancy[Beck.Nancy@epa.gov]

Thur 11/30/2017 7:04:19 PM

Dourson, Michael

```
<dourson.michael@epa.gov>; Kundinger, Kelly <kundinger.kelly@epa.gov>; Wehrum, Bill
<Wehrum.Bill@epa.gov>; Baptist, Erik <baptist.erik@epa.gov>; Lyons, Troy <lyons.troy@epa.gov>;
Darwin, Veronica <darwin.veronica@epa.gov>; Darwin, Henry <darwin.henry@epa.gov>; Brown, Byron
<brown.byron@epa.gov>; Jackson, Ryan <jackson.ryan@epa.gov>; Wagner, Kenneth
<wagner.kenneth@epa.gov>; Shimmin, Kaitlyn <shimmin.kaitlyn@epa.gov>; Kelly, Albert
<kelly.albert@epa.gov>; Forsgren, Lee <Forsgren.Lee@epa.gov>; Abboud, Michael
<abbout.michael@epa.gov>; Schwab, Justin <Schwab.Justin@epa.gov>; Rodrick, Christian
<rodrick.christian@epa.gov>; Harlow, David <harlow.david@epa.gov>; Konkus, John
<konkus.john@epa.gov>; Sands, Jeffrey <sands.jeffrey@epa.gov>; Dominguez, Alexander
<dominguez.alexander@epa.gov>; Beck, Nancy <Beck.Nancy@epa.gov>; Greenwalt, Sarah
<greenwalt.sarah@epa.gov>; Hanson, Catherine <hanson.catherine@epa.gov>; Lovell, Will (William)
<lovell.william@epa.gov>; White, Elizabeth <white.elizabeth@epa.gov>; Bennett, Tate
<Bennett.Tate@epa.gov>; Bolen, Derrick <bolen.derrick@epa.gov>; Hupp, Millan
<hupp.millan@epa.gov>; Yamada, Richard (Yujiro) <yamada.richard@epa.gov>; Bolen, Brittany
<br/><bolen.brittany@epa.gov>; Fotouhi, David <Fotouhi.David@epa.gov>; Traylor, Patrick
<traylor.patrick@epa.gov>; Bodine, Susan <bodine.susan@epa.gov>; Letendre, Daisy
<letendre.daisy@epa.gov>
> Cc: Dickerson, Aaron <dickerson.aaron@epa.gov>; Willis, Sharnett <Willis.Sharnett@epa.gov>;
Woodward, Cheryl < Woodward. Cheryl@epa.gov>
> Subject: Re: Draft LxL / No COS Meeting Tomorrow
> No COS meeting tomorrow.
> Sent from my iPhone
>
> <2016_National Analysis_Briefing_OCSPP briefinig_11-27-17.pptx>
```

To: Beck, Nancy[beck.nancy@epa.gov]

From: Dourson, Michael

Sent: Wed 11/15/2017 1:52:32 AM
Subject: RE: Glyphosate AHS publication

Nancy

Nice. But the natural question is

Ex. 5 - Deliberative Process

Ex. 5 - Deliberative Process

Cheers!

Mike

From: Beck, Nancy

Sent: Tuesday, November 14, 2017 8:46 PM

To: Dourson, Michael <dourson.michael@epa.gov>

Subject: FW: Glyphosate AHS publication

FYI

Nancy B. Beck, Ph.D., DABT Deputy Assistant Administrator, OCSPP

P: 202-564-1273
M: Ex. 6 - Personal Privacy
Beck. Nancy@epa.gov

From: Keller, Kaitlin

Sent: Saturday, November 11, 2017 12:03 PM **To:** Beck, Nancy < <u>Beck, Nancy@epa.gov</u>>

Cc: Bertrand, Charlotte < Bertrand. Charlotte@epa.gov >

Subject: Glyphosate AHS publication
Nancy—you may have already seen this but I just saw this published Thursday.
Glyphosate Use and Cancer Incidence in the Agricultural Health Study JNCI: Journal of the National Cancer Institute Oxford Academic:
https://academic.oup.com/jnci/article/doi/10.1093/jnci/djx233/4590280
Thanks,
Kaitlin
Sent from my iPhone

To: Dourson, Michael[dourson.michael@epa.gov]

From: Dourson, Michael

Sent: Thur 11/2/2017 1:43:52 AM

Subject: Doi

Note to David, note to Rick and Nancy and Jeff; note to Melissa and Chris

Sent from my iPad

To: Washington, Valerie[Washington.Valerie@epa.gov]

From: Dourson, Michael

Sent: Mon 12/18/2017 11:46:12 PM

Subject: RE: Badge

Valerie

Your badge is in your computer. Give me a call tomorrow at Ex. 6 - Personal Privacy and I will come down and check you in.

Mike

-----Original Message-----From: Washington, Valerie

Sent: Monday, December 18, 2017 6:27 PM

To: Dourson, Michael <dourson.michael@epa.gov>

Subject: Badge

Do you see my badge on my desk or in the computer?

Thanks

Sent from my iPhone

To: Beck, Nancy[beck.nancy@epa.gov]; Morris, Jeff[Morris.Jeff@epa.gov]

Cc: Bertrand, Charlotte[Bertrand.Charlotte@epa.gov]

From: Dourson, Michael

Sent: Wed 11/15/2017 1:48:08 AM

Subject: RE: Near term action on cross-federal communication - PFAS ATSDR Toxicological Profile to

be released for final public comment

Nancy and Jeff

Ex. 5 - Deliberative Process

We need that briefing from OW on their advisory pronto.

Cheers!

Michael...

... L. Dourson, PhD., DABT, FATS, FSRA

Senior Advisor to the Administrator

U.S. Environmental Protection Agency

dourson.michael@epa.gov

202-564-2463

www.epa.gov

From: Beck, Nancy

Sent: Tuesday, November 14, 2017 8:30 PM **To:** Morris, Jeff < Morris.Jeff@epa.gov>

Cc: Dourson, Michael <dourson.michael@epa.gov>; Bertrand, Charlotte

<Bertrand.Charlotte@epa.gov>

Subject: FW: Near term action on cross-federal communication - PFAS ATSDR Toxicological

Profile to be released for final public comment

Jeff,

Have we reviewed the ATSDR documents and do we feel that they have been responsive to our comments?

Thanks,

Nancy

Nancy B. Beck, Ph.D., DABT
Deputy Assistant Administrator, OCSPP
P: 202-564-1273
M Ex. 6 - Personal Privacy Beck. Nancy@epa.gov

From: Sinks, Tom

Sent: Monday, November 13, 2017 4:59 PM

To: Beck, Nancy < <u>Beck, Nancy@epa.gov</u>>; Yamada, Richard (Yujiro)

<yamada.richard@epa.gov>; Forsgren, Lee <Forsgren.Lee@epa.gov>; Grantham, Nancy
<<u>Grantham.Nancy@epa.gov</u>>; Orme-Zavaleta, Jennifer <<u>Orme-Zavaleta.Jennifer@epa.gov</u>>
Cc: Sinks, Tom <<u>Sinks.Tom@epa.gov</u>>; Rodan, Bruce <<u>rodan.bruce@epa.gov</u>>; Hubbard,
Carolyn <<u>Hubbard.Carolyn@epa.gov</u>>; Mattas-Curry, Lahne <<u>Mattas-Curry.Lahne@epa.gov</u>>;

Martin, Lawrence < Martin.Lawrence@epa.gov >; Drinkard, Andrea

<Drinkard.Andrea@epa.gov>

Subject: FW: Near term action on cross-federal communication - PFAS ATSDR Toxicological Profile to be released for final public comment

Ex. 5 - Deliberative Process

EPA is being pushed to provide communications input. While, I'm confident that EPA communications folks can provide constructive messaging related to our position and need to work on that, I don't know if additional discussions should occur beyond developing communications materials.

From: Flowers, Lynn

Sent: Monday, November 13, 2017 2:08 PM

To: Sinks, Tom <<u>Sinks.Tom@epa.gov</u>>; Martin, Lawrence <<u>Martin.Lawrence@epa.gov</u>> Cc: Hubbard, Carolyn < Hubbard. Carolyn@epa.gov>; Mattas-Curry, Lahne < Mattas-Curry.Lahne@epa.gov>; Rodan, Bruce < rodan.bruce@epa.gov>; Fleming, Megan <Fleming.Megan@epa.gov>; Burden, Susan <Burden.Susan@epa.gov>; Hauchman, Fred <hauchman.fred@epa.gov>; Sjogren, Mya <Sjogren.Mya@epa.gov>; Raffaele, Kathleen <raffaele.kathleen@epa.gov>

Subject: Near term action on cross-federal communication - PFAS ATSDR Toxicological Profile to be released for final public comment

Tom and Lawrence:

ATSDR (Henry Abadin and Janine Cory) has reached out to me to begin cross-Agency coordination on communications related to the pending release of their draft Toxicological Profile for PFAS for public comment.

The process for the document is as follows:

- 1) a non-public letter peer review (recently completed)
- 2) internal ATSDR clearance (currently ongoing)
- 3) public release for a final 60 day comment period (anticipated release in November)
- 4) final posting TBD/spring 2018

Ex. 5 - Deliberative Process

Kathleen and I would greatly appreciate being apprised of next steps as you move forward. Our PFAS toxicity workgroup is ready to help craft Qs & As.

ATSDR is expecting to hear from EPA as soon as possible.

Here is the contact information for ATSDR's communication liaison for PFAS:

Janine Cory, MPH

Associate Director for Communications (Acting)

Division of Toxicology and Human Health Sciences

Agency for Toxic Substances and Disease Registry (ATSDR/CDC)

JCory@cdc.gov

Ex. 6 - Personal Privacy

Lynn Flowers, PhD, DABT

Associate Director for Science

Office of Science Policy

US EPA

Washington, DC

202-564-6293

To: Scarano, Louis[Scarano.Louis@epa.gov]

From: Dourson, Michael

Sent: Mon 12/18/2017 11:43:18 PM

Subject: Web-link

Gino

Good to run into you. Below is a note that I sent to my University of Cincinnati colleagues giving them the news. Please feel free to share this link (<u>www.tera.org</u>, see top link on collaboration), but not the note.

Cheers!

Michael...

... L. Dourson, PhD., DABT, FATS, FSRA

Senior Advisor to the Administrator

U.S. Environmental Protection Agency

dourson.michael@epa.gov

202-564-2463

www.epa.gov

Dear Drs. Ho, Maier, and McGinnis

I have ask Mr. Trump to withdraw my name for consideration of the Assistant Administrator of EPA's Office of Chemical Safety and Pollution Prevention. This request was due entirely to the politicization of my nomination, based in large part of the misrepresentation of my prior work with the University of Cincinnati and with TERA. I sincerely apologize that the stellar work of your staffs was impugned during this process.

My withdrawal letter gives a link to the TERA website (www.tera.org, see top link on
collaboration) that gives a point by point refutation of the various mis-informed stories that
have been circulating. I appreciate Dr. McGinnis's allowing this to be placed on the TERA
website. The University should feel free to refer to this website, or create something
similar, if folks wish to see a balanced perspective.

Sincerely

Michael Dourson

To: Fugh, Justina[Fugh.Justina@epa.gov]

From: Dourson, Michael

Sent: Sat 12/30/2017 3:21:21 PM

Subject: Re: CAUTION: You're getting closer to being added to the ethics naughty list

Justina

I will be working the week of Jan 2, but this may be my last week. This, I hope you do not mind if I forgo the training. If I stay on, I will take the training. If not, then I will need to talk with you about future work.

Holiday cheers!

Michael

Sent from my iPhone

On Dec 19, 2017, at 11:06 AM, Fugh, Justina < Fugh. Justina@epa.gov > wrote:

A gentle nudge – please take your 2017 annual ethics training before 12/31/17

(and, if you read through this whole message, there's a treat at the end) ...

This year's course -- "Follow the Money – Gifts and Travel" -- examines ethics considerations related to ... you guessed it: gifts and travel! This course (the last one created by Dan Fort) will give employees an overview of the federal ethics requirements and then dive into the do's and don'ts of giving and receiving gifts, including the gift of travel. If you have questions about the course, then please contact our new training officer, Margaret Ross (whom we are delighted to have join us).

Who needs to take this training?

YOU have to take the training because you file the public financial disclosure report and, according to our records, have not yet completed training.

What is the deadline for completing the training?

Training must be completed by December 31, 2017. We know, we're tardy in reminding you, but we have faith in you!

How do you access the training?

To access the training, click on this link: <u>2017 Annual Ethics Training</u> or cut and paste the following address into your browser: http://intranet.epa.gov/ogc/2017ethicstraining/10.html. This training is still not yet hosted in Skillport; rather, it's on the OGC/ethics intranet site. If you don't have access to the intranet (sigh), then please contact Margaret Ross at ross.margaret@epa.gov.

How do you document completion of the course?

At the end of the training, you will see a screen that directs you to enter your EPA email address (i.e., lastname@epa.gov) in order to certify completion of the course. What will happen is that the system will generate a certificate in your browser, then send you an email confirming your completion, and will add your name to the tracking database in Lotus Notes. Yes, we know that Lotus Notes is going away, but OGC/Ethics will keep our subscription going into next year (and encourage ethics officials to do the same). And before our friends in OEI get upset, yes, we know that we need to move to another system next year.

What if I don't get a training certificate?

You know, if you tell me that you took the training but didn't get a certificate, well, I'll believe you. I mean, would you really lie to me about having taken *ethics* training? Just send me an email with the date you took the training, and I'll generate a certificate for you.

Why isn't the training available on Skillport?

Well, it just isn't...but it will be next year. Until then, we're going to muddle through with the Lotus Notes training tracker this last time.

How can I send feedback and comments?

As you know, our training courses are created entirely by the OGC ethics team, principally through the creative genius of Dan Fort, who recently retired. If you have any comments about the content of this course, or want to provide input about future courses, please contact Margaret Ross (our new ethics training officer) at ross.margaret@epa.gov.

PS – and now the "treat" as promised. We hear that the National Executive Leadership Development Conference (NELDC, colloquially known as the SES Forum) will be held in late January 2018. Provided you complete your online training in 2017, then you will be able to get credit for you 2018 training by attending the conference! You know that our ethics training is fun when it's in person, so please plan to stay for that!

Thanks for your attention to ethics issues! Call us anytime (or email us at ethics@epa.gov). Justina

Justina Fugh | Senior Counsel for Ethics | Office of General Counsel | US EPA | Mail Code 2311A | Room 4308 North, William Jefferson Clinton Federal Building | Washington, DC 20460 (for ground deliveries, use 20004 for the zip code) | phone 202-564-1786 | fax 202-564-1772

To: Baptist, Erik[baptist.erik@epa.gov]
Cc: Sands, Jeffrey[sands.jeffrey@epa.gov]

From: Dourson, Michael

Sent: Tue 11/14/2017 1:22:55 AM

Subject: RE: ESA

Thanks Erik!

From: Baptist, Erik

Sent: Monday, November 13, 2017 8:11 PM

To: Dourson, Michael <dourson.michael@epa.gov>

Cc: Sands, Jeffrey <sands.jeffrey@epa.gov>

Subject: Re: ESA

The letter looked good - no edits from me.

Sent from my iPhone

On Nov 13, 2017, at 5:13 PM, Dourson, Michael <<u>dourson.michael@epa.gov</u>> wrote:

Erik

How about I just meet you at Ryan's office at 6 pm? Jeff, please feel free to join us.

Michael

From: Dourson, Michael

Sent: Monday, November 13, 2017 2:56 PMTo: Baptist, Erik baptist.erik@epa.govCc: Sands, Jeffrey sands.jeffrey@epa.gov

Subject: Re: ESA

Erik

Reviewing by 5 pm is fine. I am seeing Ryan J at 6 regarding it.
Michael
Sent from my iPad
On Nov 13, 2017, at 2:43 PM, Baptist, Erik < baptist.erik@epa.gov > wrote:
Michael,
I am tied up in meetings until 5:00pm. I can review then or first thing tomorrow morning, if that works for you.
Sent from my iPhone
On Nov 13, 2017, at 2:17 PM, Dourson, Michael < dourson.michael@epa.gov > wrote
Erik
Do you have time today to look at a draft letter from DOI on the ESA issue? If so, I will walk over a copy to you.
Cheers!
Michael

To: Bertrand, Charlotte[Bertrand.Charlotte@epa.gov]; Beck, Nancy[beck.nancy@epa.gov]

From: Dourson, Michael

Sent: Mon 12/18/2017 11:04:49 PM

Subject: IRIS slides Some IRIS Slides.pptx

Charlotte



Cheers!

Michael...

... L. Dourson, PhD., DABT, FATS, FSRA

Senior Advisor to the Administrator

U.S. Environmental Protection Agency

dourson.michael@epa.gov

202-564-2463

www.epa.gov

To: Zarba, Christopher[Zarba.Christopher@epa.gov] **From:** Dourson, Michael

Sent: Thur 12/14/2017 8:27:43 PM

Subject: RE: Misc.

Chris

For balance see also www.tera.org.

Mike

From: Dourson, Michael

Sent: Thursday, December 14, 2017 1:30 PM

To: Zarba, Christopher < Zarba. Christopher@epa.gov>

Subject: Re: Misc.

Chris

Thanks for your kind words. The SAB is very important. Honeycutt is a good choice as lead.

Mike

Sent from my iPhone

On Dec 14, 2017, at 11:49 AM, Zarba, Christopher <<u>Zarba.Christopher@epa.gov</u>> wrote:

I am sorry to see you go. I was looking forward to working with you.

Best of luck in future endeavors and please stay in touch...

Christopher S. Zarba

US EPA Science Advisory Board

zarba.christopher@epa.gov

O (202) 564-0760

M Ex. 6 - Personal Privacy

To: Dourson, Michael[dourson.michael@epa.gov]

From: Dourson, Michael

Sent: Tue 11/14/2017 1:22:34 AM

Subject: RE: PFOA

C-8 FINAL CATT REPORT 8-02.pdf

Ryan

Sorry, I hit the send button too quickly. Attached is the West Virginia report. Also of note is the text on page 9.

Cheers!

Michael

2.1 Pre Meeting Action Items

TERA is a nonprofit [501(c)(3)] corporation dedicated to the best use of toxicity data for the development of risk values. This organization is very well known and respected in the toxicology arena for their professionalism, wealth of knowledge, experience, and unbiased approach to deriving risk factors. All the non-TERA toxicologists on the CATT, whether from government agencies or industry, were in unanimous support of including TERA in this project.

From: Dourson, Michael

Sent: Monday, November 13, 2017 8:17 PM **To:** Jackson, Ryan jackson.ryan@epa.gov

Subject: PFOA

Ryan

Here is the information you need for explaining the Dupont 1 ppb value (see red text below). It is from the West Virginia report in 2002. I would be more than happy to help you and Administrator Pruitt with any chemical toxicity question. I have studied most of the problematic

chemicals either while at EPA or afterwards, and sometimes both.
Cheers!
Michael
FINAL
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ASSESSMENT OF TOXICITY TEAM (CATT) REPORT
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3. 0 COMPARISON OF SCREENING LEVELS [SL] TO SITE-RELATED DATA

After the SLs for air, water, and soil were determined, DEP compared these SLs to the site-related data that has been collected to date. These comparisons are summarized below. The work of the CATT was only one facet of an investigation that continues beyond the issuance of this report. The GIST is expected to issue a report of the groundwater and surface water data in early 2003. The air modeling effort continues and is currently focusing on determining the results of the air emissions reduction efforts by DuPont required in the consent order as a 50% reduction in overall emissions (both air and water) by the end of 2003. Upgrades were completed in June 2002 which included the installation of a new scrubber and increased height of the primary C8

emissions stack.

-

<u>Water</u>

To date, of the 188 samples collected from private wells, cisterns, and springs, 50 were used for drinking water and none exceeded the 150 ppb health protective water SL for C8. Also to date, nine public water supply facilities in West Virginia have been analyzed for C8, including Belleville Locks and Dam, Blennerhassett Island, General Electric, Lubeck Public Service District (PSD), Mason County PSD, Parkersburg PSD, Racine Locks and Dam, New Haven Water Department, and Ravenswood. None of the drinking water from these facilities contained concentrations of C8 that exceeded the 150 ppb water SL. In fact, the concentrations of C8 in public water supplies were all below 2 ppb, below 15 ppb in private non-drinking water, and below 3 ppb in private drinking water wells in West Virginia. Samples were collected from Ohio public and private water supplies. Although C8 levels in some Ohio private water supplies were higher than those detected in West Virginia, none of these samples contained C8 concentrations above the water SL. These data have been provided to Ohio EPA and DEP will continue to share information with throughout the remainder of this investigation. The DEP notes that the water SL [screening level] is higher than DuPont's internal community exposure guidelines for drinking water of 1 or 3 ppb; however, these guidelines were developed in the early 1990s and based solely on a two-week inhalation study from 1986. Since then significant additional toxicological data have been collected and the CATT water SL is based on a comprehensive examination of all available information. Sampling of the Ohio River has begun; preliminary analytical results are expected from the laboratory in September 2002. To date, no analysis has been performed to measure C8 in soils in West Virginia on private property; therefore, no comparison can be made to the soil SL.

FINAL

AMMONIUM PERFLUOROOCTANOATE (C8) ASSESSMENT OF TOXICITY TEAM (CATT) REPORT



August 2002



Department of Environmental Protection - promoting a healthy environment

EXECUTIVE SUMMARY

Pursuant to a consent order signed November 14, 2001 between the West Virginia Environmental Protection and Health and Human Resources departments, and E. I. Du Pont de Nemours, Inc. (DuPont) the C8 (ammonium perfluorooctanoate) Assessment of Toxicity Team (CATT) was established to:

- (1) determine risk-based human health protective screening levels (SLs) for this unregulated chemical in air, water, and soil;
 - (2) provide health risk information to the public; and
 - (3) determine an ecological health protective SL for C8 in surface water.

To date, two public meetings have been held in the vicinity of the DuPont Washington Works facility located near Parkersburg, West Virginia. Also, a team of 10 expert toxicologists have met and determined human health provisional risk factors for the oral and inhalation routes of exposure, and calculated health protective SLs based on these risk factors using Region 9 U.S. Environmental Protection Agency standard methodology. The results of the CATT's investigation are presented in summary below. The ecological SL for surface water currently is still in development. An addendum to this report is expected to be released in Fall 2002 presenting the surface water SL findings.

The methodology, overall process, and rationale utilized by the CATT to develop these risk factors and SLs are discussed, the members are listed, and a synopsis of the events leading to the consent order are presented herein. The intent of this report is to document the process and conclusions of the CATT in an effort to provide to the public a record of these activities. It is not intended to be a summary of all the toxicology information available on C8.

The risk factor or Reference Dose (RfD) for the oral route of exposure determined by the CATT for C8 was 0.004 milligrams per kilogram of body weight per day (mg/kg-day). A risk factor for the inhalation route of exposure or the Reference Concentration (RfC) of 1 micrograms per cubic meter of air (μ g/m³) was determined. The RfD or RfC is defined by EPA as an estimate (with uncertainty spanning perhaps an order of magnitude or greater) of a daily exposure level for the human population, including sensitive subpopulations, that is likely to be without an appreciable risk of deleterious effects during a lifetime. Based on the oral RfD, health protective SLs were calculated for water of 150 parts per billion (ppb), and for soil of 240 parts per million (ppm). Based on the inhalation RfC, a health protective SL of 1 μ g/m³ was derived for air.

ACKNOWLEDGEMENTS

The West Virginia Department of Environmental Protection wishes to thank the following agencies and organizations that joined us as primary participants in this investigation: West Virginia Department of Health and Human Resources; U.S. Environmental Protection Agency (EPA) Region 3, Office of Research and Development (ORD) and Headquarters; E. I. Du Pont de Nemours, Inc. (as well as their employees, consultants - Potesta & Assoc., Inc., laboratory – Exygen Research, Inc., and attorneys); Marshall University; Toxicology Excellence for Risk Assessment (*TERA*); and Menzie Cura & Assoc., Inc. Specifically, we thank the following EPA personnel for their technical support and camaraderie: Karen Johnson, Janet Sharke, Garth Connor, Roger Reinhart, and Mary Dominiak. We also thank the following organizations for their cooperation: EPA Region 5, Ohio EPA, and the National Institute for Chemical Studies.

We thank all the individual members of the C8 Assessment of Toxicity Team (CATT) for their participation and cooperation. In particular, we thank the following CATT members:

- James Becker, M.D., and Tracy Smith, M.S., of Marshall University for their professionalism, scientific knowledge, and common sense approach to communicating environmental health risks to the public.
- The toxicologists who embarked on an expedition to find the truth, the ambition of all noble scientists:

EPA

John Cicmanec, D.V.M., M.S., USEPA ORD Samuel Rotenberg, Ph.D., USEPA Region 3 Jennifer Seed, Ph.D., USEPA Headquarters

TERA

Michael Dourson, Ph.D. Joan Dollarhide, MS, MTSC, JD Andrew Maier, Ph.D., CIH Dan Briggs, Ph.D., DABT (note taker)

Agency for Toxic Disease Registry

John Wheeler, Ph.D.

DuPont

Gerald Kennedy John Whysner, M.D., Ph.D., D.A.B.T. (consultant)

Invited guests:

John Butenhoff, Ph.D., 3M (study scientist) Jim Sferra, MS, OEPA (observer)

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1.0 INTRODUCTION

The investigation described herein was conducted pursuant to the November 14, 2001 Consent Order Number GWR-2001-019 between the West Virginia Departments of Environmental Protection (DEP) and Health and Human Resources (DHHR), and E. I. Du Pont de Nemours, Inc. (DuPont). A copy of this consent order is included as Attachment I. These actions were instigated by the presence of an unregulated chemical, ammonium perfluorooctanoate commonly called C8, in the Lubeck, W.Va. public water supply which is near the DuPont Washington Works (WW) facility in Washington, W.Va. A site map is included in Attachment IIc.

The consent order established two scientific teams: (1) the C8 Assessment of Toxicity Team (CATT), and (2) the Groundwater Investigation Steering Team (GIST). The CATT was tasked with investigating the toxicity of C8; developing provisional risk factors for the inhalation, dermal, and oral routes of exposure; and establishing human health protective screening levels (SLs) for air, water, and soil; investigating the ecological toxicity of C8 and determining an ecological health protective SL for surface water; and with communicating health risk information to the public. In the consent order DuPont agreed to meet these SLs at their WW facility, once developed, and that these SLs would remain in effect until superseded by U.S. Environmental Protection Agency (EPA) guidance. The CATT's activities and findings regarding the toxicity of C8, development of risk factors and SLs are presented in detail in Section 2 of this report. Slides presented at the two public meetings held thus far are provided in Attachment II. The investigation into the ecological toxicity of C8 and surface water SL development is scheduled for completion in Fall 2002. When finished, the surface water will be presented in an addendum to this report.

The GIST was established by the consent order to determine the extent and concentration of C8 in both groundwater and surface water. The activities of the GIST continue as of the issuance of this CATT report. The GIST will issue a report on the C8 analytical data for groundwater and surface water when that work is finished, scheduled for early 2003. Interim reports are available through the DEP Division of Water Resources (DWR). The groundwater investigation focused not only on the WW plant, but also on areas where C8 had been disposed, including the Local Landfill (on WW property), Dry Run Landfill (near the WW plant), and the Letart Landfill (30 miles south of the WW plant). Maps of the one-mile radius study area around these locations are included in the presentation of interim results at the second public meeting provided in Attachment IIc.

Summarized findings to date by the GIST are compared to the health protective water SL developed by the CATT in Section 3.0. Results of air dispersion modeling efforts thus far conducted by the DEP Division of Air Quality (DAQ) are compared to the air SL in Section 3.0 as well.

Background

The DuPont WW plant is located approximately 10 miles southwest of Parkersburg, W.Va. along state Route 61 in the rural hamlet of Washington, W.Va. This facility was established in the 1940s and currently is one of the largest DuPont enclaves in the world. DuPont has used C8 at this facility for more than 50 years as a surfactant in various manufacturing processes, including the production of Teflon. "C8" is the 3M trade name for its product that contains ammonium perfluorooctanoate (APFO) (CAS # 3825-26-1). In biologic media, APFO quickly dissociates to perfluorooctanoate, which is the anion of perfluorooctanoic acid (PFOA). The PFOA form has been identified as potentially toxic to animals. Throughout this report, C8 is used as terminology to include C8, APFO, or PFOA.

The DEP became aware of and began investigating the presence of C8 in the Lubeck, W.Va. public water supply in November 2000. In Spring 2001, DEP received a letter requesting a formal agency investigation into DuPont's environmental releases of C8 and the presence of C8 in the Lubeck drinking water from attorneys representing a few citizens residing in proximity to the WW plant. The Lubeck public water supply well field lies approximately 3 miles south of the DuPont WW plant. Also around this time, DEP became aware that C8 was chemically similar to perfluorocate sulfonate (PFOS), another perfluorocarbon manufactured by 3M, and that 3M had recently removed their Scotchguard product from the marketplace because it contained PFOS. From U.S. EPA Region 3 and Headquarters, DEP learned that 3M had undertaken a significant research effort into the toxicity of perfluorocarbons, particularly PFOS and including C8; that perfluorocarbons were potentially more toxic than previously thought; that 3M was submitting the new data to EPA under the Toxic Substances Control Act (TSCA); and that these data were publicly available under Administrative Record 226 (AR226). Additionally, DEP learned that DuPont was submitting toxicity data on C8 to EPA, as well.

DEP gathered data and met with DuPont and met with citizens attorneys in Spring 2001. The DEP, which regulates groundwater in West Virginia, was joined in the investigation by the DHHR, which regulates drinking water. The DHHR requested support from EPA Region 3 to enforce the National Safe Drinking Water Act. At the request of these agencies, DuPont supplied information regarding C8 and its use in manufacturing processes, its toxicity, and emissions. After several months of investigation and discussions, a consent order was signed in November 2001. A copy of the consent order is provided in Attachment I. It describes the tasks and members of the CATT and GIST. The DEP informed the public of the consent order and scheduled a public meeting to discuss the order.

The DEP held it's first public meeting regarding C8 on November 29, 2001 at Blennerhassett Junior High School which is located near the Lubeck and Washington communities. The meeting was spearheaded by the CATT and the GIST. The purpose of the meeting was to inform citizens of: (1) the requirements of the consent order; (2) the members and activities of the GIST; (3) their assistance was required to fill out and return a water use survey if they had groundwater wells, cisterns, or springs (particularly those used for drinking water), and to allow sampling of these water sources; (4) the members and activities of the CATT; (5) the available information regarding the toxicity of C8; and (6) the known current levels of C8 in the Lubeck public water supply, which were below 1 part per billion (ppb). At this meeting, James Becker, M.D. of Marshall University spoke regarding environmental exposures and risks in general, and Dee Ann Staats, Ph.D. (DEP) explained the CATT and GIST activities, the consent order, and known toxicity of C8. The slides from both presentations are provided in Attachment IIa.

By the end of January 2002, contractors were in place to assist the CATT and the GIST in their tasks. The GIST was headed by DEP and had members from DHHR, EPA, and Dupont. The CATT was headed by DEP and had members from DHHR, EPA, DuPont and the Agency for Toxic Substances Disease Registry (ATSDR). The DEP contracted with the National Institute of Chemical Studies (NICS), a nonprofit organization, which subcontracted the human and ecological toxicology work to the Toxicology for Excellence in Risk Assessment (*TERA*) group, also a nonprofit, which subcontracted the ecological toxicology work to Menzie Cura & Assoc., Inc. (MC). Both *TERA* and MC are well respected in the field of toxicology. The NICS subcontracted the risk communications tasks to Marshall University.

In March 2002, EPA Regions 3 and 5 signed a consent order with DuPont requiring the provision of alternative water to any resident in West Virginia or Ohio with C8 in drinking water at levels above 14

ppb. The 14 ppb was an interim value in effect until the water SL was developed by the CATT. This value was taken from the final report by ENVIRON Int. Corp. (a consulting firm hired by DuPont) titled "A Hazard Narrative for Perfluorooctanoate (PFOA)", January 2002. An earlier draft, "A Review of the Toxicology of Perfluorooctanoate (PFOA)", November 2001, had proposed a drinking water value of 210 ppb. However, DEP's toxicologist, Dr. Staats, expressed concern over some of the assumptions made in the calculation of the 210 ppb to DHHR and EPA Region 3. The outcome of these discussions was a decision that a very conservative approach should be taken in the interim until the CATT water SL was developed. Therefore, 14 ppb was accepted as the interim water SL for alternative water provision. Note that this consent order was jointly signed by two regions of EPA because West Virginia is in Region 3 and Ohio is in Region 5. During the investigation, C8 had been found in the Little Hocking, Ohio public water supply. Also, note that DEP and DHHR invited Ohio EPA to join the CATT and GIST as observers, but not as members because this would have required renegotiating the consent order between West Virginia and DuPont.

TERA was assigned by DEP to review and compile the C8 toxicological information provided by DEP and to prepare for and hold a meeting of the CATT toxicologists during which the provisional risk factors and health protective SLs would be derived. The CATT toxicologists panel was comprised of 10 expert scientists with a collective span of experience of over 175 years and many specialties including endocrinology, veterinary medicine, cancer, and risk assessment.

TERA's efforts are described further in Section 2.1. By mid April 2002, TERA was prepared for the meeting. Also, TERA helped prepare the other toxicologists for the meeting by providing toxicity reports and summary information. The CATT toxicologists met on May 6 and 7, 2002 at EPA offices in Cincinnati, Ohio. The minutes of this meeting are provided in Section 2.2. The meeting lasted approximately 18 hours with roughly one-third of that time spent in discussions of C8's potential carcinogenicity. The oral provisional reference dose (pRfD) risk factor, and the two health protective SLs (for water and soil) based on this risk factor were developed at this meeting. The panel agreed that the toxicology database was insufficient to develop a dermal exposure pRfD. The inhalation provisional reference concentration (pRfC) risk factor and air SL developed at the meeting were only interim because additional data collection was necessary for their calculation. These data were collected and provided to TERA, who calculated the final pRfC and air SL, wrote a report describing this activity and forwarded it to the other CATT toxicologists for their approval. This document is provided in Section 2.3 as the post meeting action items. Both the meeting minutes and the post meeting action items were reviewed and approved by the panel of 10 highly qualified toxicologists.

An internal briefing for the DEP, DHHR, and EPA was held on May 8, 2002 to discuss the water and soil SLs. Rather than withhold this information while the meeting minutes report was prepared, DEP released the water and soil SLs so that the public would be informed of the status of their drinking water, and decisions could be made regarding the provision of alternative water supplies. In that spirit, DuPont and the public were informed – via a meeting with the above regulators and a press release, respectively - of the water and soil SLs on May 9, 2002.

A second public meeting was held at Blennerhassett Junior High School on May 15, 2002, to inform the public of the details of the SL development and of the groundwater C8 concentrations that had been detected at that point. Dr. Becker first spoke regarding environmental health risks in general. Dr. Staats described the process used by the CATT toxicologists to arrive at the water and soil SLs. Finally, David Watkins (DEP, GIST chairman) presented the C8 analytical data for private and public water sources. Slides of the presentations given at this meeting are provided in Attachment IIb.

2.0 DEVELOPMENT OF RISK FACTORS AND SCREENING LEVELS

TERA was assigned to prepare for, host and document the meeting of the CATT toxicologists during which the provisional C8 risk factors (pRfDs and pRfC) would be developed by the group. The activities undertaken by TERA to prepare for the meeting are presented in Section 2.1. The actual minutes of the meeting are provided in Section 2.2., and the tasks conducted by TERA to develop the final air SL after the meeting at the direction of the panel are described in Section 2.3.

2.1 Pre Meeting Action Items

TERA is a nonprofit [501(c)(3)] corporation dedicated to the best use of toxicity data for the development of risk values. This organization is very well known and respected in the toxicology arena for their professionalism, wealth of knowledge, experience, and unbiased approach to deriving risk factors. All the non-TERA toxicologists on the CATT, whether from government agencies or industry, were in unanimous support of including TERA in this project.

TERA was tasked with compiling and reviewing the available toxicological data for C8. A literature search and review of these data was in draft by EPA Headquarters, this document was provided to TERA. The 3M submittals to AR-226 were provided to TERA by DEP. These data grew from a total of seven compact discs to 10 during the time period of this project. The AR-226 continues to grow with 3M submittals currently. The index of the first seven discs are provided in Attachment Va. Additionally, DEP conducted a literature search of C8 toxicity data on the National Library of Medicine's Medline and Toxline databases in June 2001. The results of these searches were provided to TERA by DEP as well. Also, documents submitted to DEP from DuPont in response to the EPA Region 3 request for information was made available to TERA by DEP, first by mailing relevant toxicology documents identified by Dr. Staats, and then by physically delivering all these documents to their Cincinnati office for TERA to sort and identify those deemed relevant and necessary for their work. Therefore, little literature searching or data retrieval was required of TERA.

After reviewing the existing C8 toxicology data, TERA selected studies that would be suitable for derivation of risk factors for the oral, dermal, and inhalation route of exposure. A list of the potential key studies was prepared. An indepth review of these studies was then conducted, and the details of the studies were summarized in tabular format. Next, TERA prepared a condensed table of these studies including critical effects and exposure levels identified by TERA, and blank columns for the other criteria necessary in the risk factor development process, such as the uncertainty factors. The documents listed below were provided to the other CATT toxicologists approximately two or three weeks prior to the meeting. TERA also prepared tables of suggested uncertainty factors, risk factors, and resulting SLs to DEP. These documents were discussed with Dr. Staats but were not distributed to the other toxicologists prior to the meeting in an effort not to influence their decisions, and not to give the false impression that the decisions on risk factor development had already been made and that the panel's purpose was simply to review TERA's work. Rather, TERA's suggestions would be presented at the meeting as a starting point for panel discussions and the development of the risk factors and SLs would be done as a group. The pre-meeting documents provided to the rest of the panel by TERA and DEP are contained in Attachment III. Also in Attachment III is a more detailed description of the decisions and methodology used by TERA in suggested risk factor development.

2.2 CATT TOXICOLOGISTS MEETING MINUTES

Meeting of C8 Assessment of Toxicity Team (CATT) Toxicologists

May 6 and 7, 2002

Andrew W. Breidenbach Environmental Research Center, Cincinnati, Ohio

Attendees:

Voting Team Members

John Cicmanec, D.V.M., M.S., ACLAM, USEPA Office of Research and Development Joan Dollarhide, M.S., M.T.S.C., J.D., Toxicology Excellence for Risk Assessment (*TERA*) Michael Dourson, Ph.D., D.A.B.T., *TERA* Gerald Kennedy, E. I. Du Pont de Nemours, Inc. Andrew Maier, Ph.D., C.I.H., *TERA* Samuel Rotenberg, Ph.D., USEPA Region 3 Jennifer Seed, Ph.D., USEPA Office of Pollution Prevention and Toxics (may abstain from voting) Dee Ann Staats, Ph.D. (Chairperson), West Virginia Department Environmental Protection (DEP) John Wheeler, Ph.D., D.A.B.T., Agency for Toxic Substances Disease Registry (ATSDR) (representing West Virginia Department of Health and Human Resources [DHHR]) John Whysner, M.D., Ph.D., D.A.B.T. (consulting for DuPont)

Invited Guests

John Butenhoff, Ph.D., 3M Company (study director) Jim Sferra, M.S., Ohio EPA (observer)

Note taker

Daniel Briggs, Ph.D., D.A.B.T., TERA

Introduction

The toxicologists on the C8 Assessment of Toxicity Team (CATT) met on May 6 and 7, 2002, to develop provisional reference doses (pRfDs) and screening levels (SLs) for ammonium perfluorooctanoate (C8) as specified in Consent Order GWR-2001-019 between the West Virginia Department of Environmental Protection, the West Virginia Department of Health and Human Resources, and E. I. Du Pont de Nemours & Co., (DuPont) dated November 14, 2001. These screening levels apply only to DuPont at their West Virginia facilities as specified in this consent order. Any use of these pRfDs or SLs for any other purpose or by any other regulatory agency is solely their choice and responsibility.

The meeting opened with Dr. Staats announcing that this meeting was being held pursuant to the above-cited consent order as part of an enforcement action and was therefore closed to the public. Dr. Staats noted that, except for Dr. Butenhoff and Mr. Sferra who were invited guests, the panelists were named as part of the consent order and were free to enter into discussions and vote on issues. It was noted that Dr. Seed could abstain from voting at any time. The rules for the meeting were set forth as follows:

- The panel would strive for unanimous consensus, but if such consensus could not be reached, then the majority of votes would rule.
- The panel was expected to be cooperative and courteous with each other.
- The risk factors and screening levels would be developed together as a group, rather than simply by reviewing the work and suggestions of *TERA*.
- Votes would be taken at each decision point. After panel discussion on each point, a
 motion would be made on the floor. The chair would then repeat the motion and
 verbally poll each panel member individually. The chair would always vote last in
 order to not influence the voting.

TERA recorded the official minutes for the meeting. However, the chair recorded supplemental notes, which were provided TERA to assist in the preparation of the final Meeting Minutes Report. It was noted that specific discussion comments or votes would not be attributed to panel members (i.e., no names would be used) in the meeting report in order to facilitate full and open discussion among the team. It was also noted that TERA would distribute a draft meeting report to the CATT panel for their review and incorporate panel comments as appropriate. Each panel member would be asked to sign a statement agreeing that the meeting report is an accurate representation of the discussion and conclusions of the CATT Team. The original signatures will remain on file with the DEP.

The sequence of discussion on Monday, May 6 was oral noncancer assessment; dermal noncancer assessment and on Tuesday, May 7 was cancer assessment; inhalation noncancer assessment; oral screening level; and interim inhalation screening levels. (Note that Dr. Seed left the meeting at 2:30 pm on Tuesday, May 7, 2002; she was present and joined in all discussions through the cancer assessment.) However, for clarity, the meeting report is organized according to noncancer (oral, dermal, inhalation) assessment, cancer assessment, and screening levels. Below, under each heading is a brief description of *TERA*'s opening comments, followed by the panel discussion, and then the outcome of the panel discussion.

Noncancer Assessment: Review of the Oral Studies

Prior to the meeting, *TERA* evaluated the available human and animal health effects studies for C8. (A list of the documents and studies included in *TERA*'s prior review is provided in the Attachments). *TERA* evaluated the pool of available studies to identify the key studies that could be selected by the CATT panel as the basis for the pRfD. In narrowing the list of available studies, the available data were evaluated weighing considerations such as observed effect levels, study duration and quality, and applicability to human health. The judgments were made in a manner consistent with hazard identification and dose-response assessment practices used in current U.S. EPA risk assessments. Studies were generally given greater consideration as potential principal studies if they were at least of subchronic duration; identified NOAEL/LOAEL boundaries on the low end of the range provided by all the data; and had robust design (e.g., diverse array of endpoints, sufficient number of animals). From the total pool of available studies, *TERA* developed detailed summary tables for each of the key

studies having potential for being selected as the principal study for derivation of the pRfD. The resulting detailed summary table of key studies was provided to the panel members prior to the meeting to facilitate the selection of the principal study by the CATT panel and is attached. Therefore, discussion of the oral studies at the meeting focused on the tables presented in the attachment which identified those studies of sufficient duration, content, and quality to merit consideration as the bases for deriving a pRfD. The tables present *TERA*'s selection of critical effect levels, and highlight the study data for key parameters that showed treatment-related changes.

At the opening of the meeting, the panel discussed whether all adequate studies had been included and whether any potential key studies were missing. One panelist asked why the 90-day Rhesus monkey study (Goldenthal, 1978b) had not been included. *TERA* responded that the Rhesus study was not considered to be as useful as the cynomolgus monkey study (Thomford et al., 2001) because it had fewer animals per group, and suggested a higher NOAEL/LOAEL boundary; however, findings from the Rhesus study would be discussed together with the cynomolgus study as supporting data. The panel confirmed that, to the best of their knowledge, the table included all of the toxicity work that should be considered in selecting principal studies for deriving the pRfD for C8.

After agreeing that all of the potential critical studies had been identified, the panel then discussed the merits of each of the studies, and the appropriate No-Observed-Adverse-Effect-Levels (NOAELs), Lowest-Observed-Adverse-Effect-Levels (LOAELs), and lower bounds on the benchmark doses (BMDLs) for each study.

<u>Human Studies (Olsen et al. 2000; Olsen et al. 1998; Gilliland and Mandel 1996; Gilliland and Mandel 1993; Ubel et al. 1980)</u>

TERA initiated the discussion by providing a brief synopsis on the potential utility of the available human health effects studies for deriving the pRfD. Two cohort mortality studies were available: (1) Ubel et al. (1980) reviewed the records of 180 deceased 3M employees for a period of 30 years (1948-1978) and found no significant difference between observed and expected mortality rates; (2) Gilliland and Mandel (1993) found no increases in mortality rates from liver cancer or liver disease in 3,537 (2,788 males and 749 females) exposed 3M workers for 35 years (1947 – 1983). Note that since the CATT meeting, a new epidemiological study on almost 4,000 (80% male) 3M workers has been completed which found no increase incidence of cancer in C8 exposed workers. Several crosssectional studies of 3M workers (111, 80, and 74 males in 1993, 1995, and 1997, respectively) were available. However, these studies were noted as being limited for use in deriving the pRfD, since workers were exposed to unknown amounts of C8 for varying time periods, and no clear signs of toxicity (such as elevated serum levels of liver enzymes were reported). The mixed findings regarding changes in hormone levels were noted. It was noted that many of these studies provided data on serum levels of C8 (or serum fluorine levels), which could serve as a measure of exposure. However, the current toxicokinetics data were not viewed as sufficiently developed to conduct a quantitative extrapolation from the reported serum levels to equivalent oral doses in humans. Based on this introduction, the panelists were asked to comment on the human data and its usefulness for deriving the pRfD.

<u>Key Panel Discussion Points:</u> Panelists noted that, although limited, the existing human data are consistent with the animal data when exposure levels are considered. Although weaknesses in the epidemiology data were noted, one panel member commented that the human data are useful for hazard identification purposes, and provide some level of comfort in conducting the assessment since they do not identify adverse effects in chronically exposed workers. It was noted that a few of the

human subjects had C8 serum levels comparable to those observed in animal studies [20 parts per million (ppm) or greater]. Other panel members described gaps in the human studies. Regarding the absence of effects observed in the epidemiology studies, the panel noted that the small number of female subjects and uncertainties in exposure levels for workers prevents the existing data from being used to rule out human toxicity. For example, the very small numbers of women in the studies prevent drawing a conclusion regarding female reproductive effects. One panelist noted that the increased blood level of estradiol reported in some subjects is not clinically significant. In addition, no adjustments were made for body mass index (BMI) variations among subjects. Since BMI is known to affect estradiol levels and in this study BMI was the only parameter to correlate with hormone levels, it was noted that it is unlikely that C8 exposure was related to increased estradiol levels. The panel discussed Gilliland and Mandel (1986), which reported six prostate cancer deaths overall and four among exposed workers. One panel member commented on the update to this study (no study report was provided), which showed no indication of increased risk of prostate cancer. This follow up study demonstrated that only one of the four workers with prostrate cancer were determined to have been exposed when work history records and blood levels of C8 were examined.

It was suggested that it might be possible to correlate C8 serum concentrations with lack of observed toxicity to estimate a human NOAEL. However, it was noted that the lack of clear exposure levels in the human studies precluded this type of analysis. Although C8 half-life determinations were conducted in some of the human studies, this information cannot be used to determine exposure doses because some exposure to the subjects may still be occurring. However, it is clear that humans do not have the major sex-related half-life difference that exists in rats. It was noted that a physiologically-based pharmacokinetic (PBPK) model is being developed, which may be useful in estimating exposure concentrations from human serum C8 levels. However, a panel member familiar with the status of this current toxicokinetic modeling effort, noted that the data are not sufficiently developed to use for quantitative risk assessment purposes at this time.

Outcome: The panel agreed unanimously that the human studies were not adequate to be used for quantitative dose-response determinations. The human studies have many substantial data gaps, such as low numbers of subjects and unknown exposure concentrations. No LOAEL was established and the exposure uncertainty does not allow identification of a clear NOAEL. In final comments made during polling of the panel, one panel member agreed with the group, but noted that the data could be used to develop a bounding estimate. A second panel member added that some evidence suggests the endocrine system as a target for C8 effects, and therefore, the human data might support the animal toxicity studies.

Definition of Adverse Liver Effect

TERA noted that in all experimental animal studies liver effects occurred. For the purposes of conducting this assessment, TERA defined adverse liver effects as the presence of histopathology (moderate grade hypertrophy would be considered sufficient) in addition to statistically significant absolute or relative weight changes, or a liver weight change of 10% or greater. A doubling of serum levels of liver enzyme activity (e.g., alkaline phosphatase (ALP), aspartate aminotransferase (AST), or alanine aminotransferase (ALT)) would also indicate an adverse liver effect. These adverse effects are used by other health organizations as well. The panel unanimously agreed with this general definition of adverse for liver effects, but noted that individual studies could demonstrate a continuum of liver effects that could be considered biologically significant.

Palazzolo et al. 1993

This is a 90-day study in male rats in which animals received C8 at doses of 0, 0.05, 0.47, 1.44, and 4.97 mg/kg-day in feed. The major finding in this study was increased liver weight with histopathological findings such as moderate hypertrophy. Panelists were asked to comment on the data from this study; on the selection of study adverse effect levels; and on the usefulness of this study as the basis for deriving a pRfD.

Key Panel Discussion Points: The possible role of peroxisome proliferation in the observed liver effects was discussed. The panel discussed uncertainty in the relevance of this mechanism to humans. One panelist stated that when considering the relevance of peroxisome proliferation, it is important to consider both qualitative and quantitative issues. This panelist suggested that peroxisome proliferation may potentially occur in humans because the cellular receptor that modulates this reaction in rodents has been found in humans, but that this mode of action should be considered to be only qualitatively relevant to humans because the receptor is far less expressed in humans, and humans have not been shown to manifest a peroxisome proliferation response. It was noted that USEPA has an ongoing project to investigate the relevance to humans of rodent peroxisome proliferation effects, but at this time EPA has no official policy on the significance of peroxisome proliferation for humans. It was also noted that IARC has also considered the issue of peroxisome proliferation and concluded that this mode of action is not relevant to humans if it has not been demonstrated to occur in human cells or primates treated with the chemical in question. (Note that the panel discussed the role of peroxisome proliferation as a potential mode of action for tumor formation later in the meeting. The results of this discussion are documented in the section on Cancer Mode of Action)

Discussion occurred regarding the usefulness of relative versus absolute liver weight in determining adverse effect levels. One panelist stated that changes in both of these parameters are preferred before designating a dose as an adverse effect level. However, most panelists considered a change in relative liver weight to be sufficient to designate a dose level as an adverse effect level. It was noted that liver weights in dosed animals in this study were comparable to control values after an 8-week recovery period; however, the panel agreed that this recovery should not influence selection of the NOAEL and LOAEL values.

Outcome: The panel agreed unanimously that 1.44 mg/kg-day is the LOAEL for this study because at this level statistically-significant increases in relative liver weight and CoA oxidase activity occur. In addition, hepatocellular hypertrophy of minimal severity or greater is observed in 14 of 15 animals at this dose, and in 2 of 15 animals at grade 2 or higher. The panel recommended that benchmark dose modeling be performed for the data based on grade 2 or higher hepatocyte hypertrophy. This modeling was conducted during the course of the meeting, resulting in a BMDL estimate of 1.3 mg/kg-day. It was noted that this BMDL is essentially the same as the LOAEL found in this study. Most panelists believed 0.47 mg/kg-day is the NOAEL because at this dose there are no statistically significant changes in either absolute or relative liver weight and only a "minimal" severity of hepatocellular hypertrophy is reported at this dose. However, one panel member preferred to call this a "minimal LOAEL" rather than a NOAEL, noting that dose-related changes in critical liver parameters had been established at the lower dose levels and suggesting that these could be part of the continuum of effects that might be considered a minimal LOAEL.

Goldenthal 1978a

This is a 90-day study in male and female rats in which animals received C8 in their feed at doses of 0, 0.56, 1.72, 5.64, 17.9, or 63.5 mg/kg-day for males and 0, 0.74, 2.3, 7.7, 22.4, or 76.5 mg/kg-day for females. This study is limited by the small number of animals (5/sex) in each dose group. Therefore, this study was not considered to be a key study. However, it was presented for the panel's consideration and comments because it includes female as well as male animals and the data on relative liver weights allow a BMD to be calculated.

<u>Key Panel Discussion Points:</u> One panelist noted that a sex difference was observed in this study. Another mentioned that this study demonstrates the importance of internal dose (C8 serum level), as compared to the administered dose.

<u>Outcome:</u> The panel agreed with the proposed NOAEL, LOAEL, and BMDL as presented by *TERA*. However, the panel also agreed unanimously that the study was not adequate to serve as the basis for deriving a pRfD because of limitations in the study (e.g., the small number of animals).

York 2002

This is a two-generation reproduction study in which male and female rats received C8 doses of 0, 1, 3, 10, and 30 mg/kg-day by gavage in distilled water. Parental animals were exposed through cohabitation and gestation to weaning of F1 animals, approximately 6 weeks. F1 animals were exposed from weaning until weaning of the F2 generation. The primary findings were increased liver weight and liver pathology in P and F1 generation male animals; however, it was noted that histology was conducted only when gross effects had been observed, and therefore liver histopathology data were not available for the control and low-dose F1 generation males.

Key Panel Discussion Points: One panelist stated that this was study was of excellent quality because it was conducted according to OPPTS guidelines for 2-generation studies. Two panelists noted that the degree of F1 generation exposure to C8 while in utero and while nursing was uncertain and may not have occurred at all because of rapid elimination of C8 from the systemic circulation of the female rats after it was administered via gavage. Therefore, the lack of reproductive toxicity in this study may not be meaningful. Other panelists agreed, but stated that the fact of rapid clearance resulting in decreased fetal exposure may not be relevant for humans because women do not have the same active secretory mechanism for C8 that exists in the female rat. Another panelist noted that rodent placenta provides less of an anatomical barrier than exists in primates. Another panelist observed that studies with radiolabeled C8 demonstrated that C8 could cross the placental barrier in rats. One panelist wondered whether female rat pups at weaning have developed the active secretory mechanism for C8 that exists in the mature females. Another panelist recalled data showing that weanling female rats were able to clear C8 faster than males, but not as fast as mature females. One panelist recommended that delayed sexual maturation and increased frequency of estrous cycles be included in the adverse effects noted for females for this study. A panelist pointed out that this study indicated a critical difference in the toxicity of C8 versus the structurally similar perfluorocarbon PFOS; in that PFOS caused fetal death at birth in a similarly designed study, while in this study C8 administration was associated with only a slightly statistically significant increase in fetal death at the post-weaning timeframe.

<u>Outcome</u>: The panel concluded that the LOAEL for males is 1 mg/kg-day. The males showed statistically-significant increases in liver and kidney weights at 1 mg/kg-day. No histology was conducted on liver and kidney at this dose level because no gross lesions were seen. However, given

the substantial histopathology noted at the next higher dose level (3 mg/kg-day), the panel believed pathology does exist at the 1 mg/kg-day level; therefore this level meets the agreed-upon definition of an adverse effect. The panel concluded that the LOAEL for females is 30 mg/kg-day. The females showed several adverse effects at this dose level, including increased mortality and decreased body weight. No NOAEL was identified for males; the NOAEL for females is 10 mg/kg-day. All of these values apply to both the P and F1 generation animals. Two panel members reviewed the BMDL modeling results, and agreed with the selection of 0.42 mg/kg-day as the study BMDL.

Riker Laboratories 1983

This is a chronic, 2-year study in male and female rats in which animals received C8 in feed at doses of 0, 1.3, and 14 mg/kg-day for males and 0, 1.6, and 16 mg/kg-day for females. The primary findings in this study are liver effects in male rats. However, it was noted that this chronic study also reported non-hepatic effects (ovarian stromal hyperplasia and ataxia) in female rats. Although this effect was not found in the subchronic study that included females (Goldenthal, 1978), the small number of animals in that subchronic study (n=5) may have limited the power of the study to observe these effects.

Key Panel Discussion Points: One of the panelists identified some copying errors in the tables (incidences of mammary fibroadenomas, Leydig cell adenomas, and ALT activity in the control group) and these values were corrected prior to the panel discussion (the attached table presents the corrected values). The panel disagreed with the study author's conclusion stated in the study report that the testicular vascular mineralization was a "spontaneous change occurring in aging rats" and that the ovarian stromal tubular hyperplasia was "equivocally related" to C8 administration because it did not progress. The panel considered both these effects to be biologically significant and relevant for determining adverse effect levels. One panelist stated that ovarian stromal hyperplasia is not commonly found in rats and noted that in this study the incidence of ovarian stromal hyperplasia in the control animals is zero. The panel discussed the relevance of the ataxia observed in females, but did not reach any conclusions about its possible biological significance. One panelist noted that at the time this study was conducted, the term "hepatic megalocytosis" was synonymous with the term "hepatic hypertrophy" currently in use. It was noted that the BMDL of 0.73 mg/kg-day calculated based on liver effects in males is consistent with the NOAELs for liver effects observed in other rat studies. In the initial summary table from which the panel was working it was noted that no BMDL was estimated for ovarian stromal tubular hyperplasia, since an adequate fit to the data was not achieved. One reviewer suggested that a model fit might be possible using log-transformed data, since the study results showed a clear log-related response curve. This approach was applied during the meeting, and resulted in a best estimate of the BMDL of 1.6 mg/kg/day.

Outcome: The panel agreed unanimously to the proposed NOAEL of 1.3 mg/kg-day for males, with a corresponding LOAEL of 14 mg/kg-day based on the following adverse effects: increased liver weight, hepatic cystoid degeneration, increased ALT enzyme activity, and testicular vascular mineralization. The panel agreed that the LOAEL in females was 1.6 mg/kg-day based on a statistically significant increase in the incidence of ovarian stromal tubular hyperplasia, and that this study did not identify a NOAEL for females. The panel further agreed that the estimated BMDL from this study is 0.73 mg/kg-day based on liver effects in males as the benchmark response.

Thomford et al., 2001

This is a 26-week study in cynomolgus monkeys, in which animals received C8 at doses of 0, 3, 10, or 30/20 mg/kg-day by gastric intubation of gelatin capsule. Gastric capsule intubation was chosen as the method of C8 administration to avoid emesis, which had occurred in the earlier Rhesus monkey study (Goldenthal et al., 1978b). Even so, several animals had problems tolerating the highest C8 dosing; as a result, the high dose was either reduced or in some cases, discontinued. Afterwards, time-weighted average doses were used to approximate the C8 dose given to the high-dose group. One animal died in the high dose group; primary findings included clinical signs and altered liver weight. TERA presented that altered liver weight was not considered an adverse finding.

Key Panel Discussion Points: At least two panelists believed that the degree of absolute liver weight increase (30%) noted at the 3 mg/kg-day dose should be sufficient to identify this dose as the LOAEL. Other panelists responded that this weight increase resulted from mitochondrial proliferation, and therefore was an adaptive response, not an adverse effect. They also pointed out that, unlike laboratory rodents, cynomolgus monkeys routinely exhibit large genetic variations. As a result, large differences in organ weights among these animals is relatively common and a 30% difference between groups – especially small groups, as in this study – is not necessarily biologically meaningful. Some panelists attempted to compare this study with the study conducted in Rhesus monkeys in order to help define the LOAEL, but this was not possible due to the uncertainty of dosing caused by the emesis that occurred in the Rhesus study. One panelist asked if the dosing technique (gastric intubation of the drug contained in gelatin capsules) might have contributed to a large range of C8 blood levels because of differences in capsule disintegration rates. Another panelist responded that this was unlikely because, while the data sometimes demonstrated large inter-animal variations in blood levels, the intraanimal variation over several dose administrations was small. It was noted that C8 serum levels were essentially the same in the low and mid-dose groups: 74, 80, and 120 µg/mL at 3, 10, and 30/20 mg/kg-day, respectively. The panel concluded that the similarities in serum C8 levels may explain the very similar effects observed between the 3 and 10 mg/kg-day dose groups. One panelist noted that protein-binding saturation was similar between the monkey and human.

<u>Outcome</u>: The panel agreed that the LOAEL is best described as "from 3 to 10 mg/kg-day" based on 30% increased absolute liver weight, and that a NOAEL does not exist for this study. At all three dose levels, statistically significant increases in absolute and relative liver weights occurred, but without accompanying histopathology. No clinical or histopathological evidence of organ damage occurred at any of the three dose levels. Dose-related trends toward lower T3 and T4 levels were observed, but these failed to achieve statistical significance, even at the highest dose. The panel concluded that these data are insufficient to identify any <u>single</u> dose as a LOAEL or NOAEL. Since the serum C8 levels were essentially the same for both the 3 and 10 mg/kg-day doses, the panel believed that designating a range of 3 to 10 mg/kg-day for the LOAEL is the best way to describe the study results.

Noncancer Assessment: Oral Hazard and Dose-Response Characterization

(Note: Dr. Seed abstained from voting during this part of the meeting.)

Critical Study and Point-of-Departure

The summary of NOAELs, LOAELs, and BMDLs unanimously agreed to by the panel is presented in Table 1 below. The individual study adverse effect levels were discussed by the panel for the purpose of selecting a critical study and effect level for derivation of the pRfD.

Key Panel Discussion Points: The primary target organ for C8 is the liver, and males are clearly more sensitive to this effect than female rats. One panelist observed that the liver effects in rats may be related to peroxisome proliferation, and therefore may not be quantitatively relevant for humans. For this reason, the liver effects in rats might not be an appropriate critical endpoint. Another panelist responded that, because of this, it was important to note that the monkey and rat LOAELs are in the same range, and since the liver effects in monkeys may not be related to peroxisome proliferation, liver toxicity might also be a relevant endpoint for humans. The observation of ovarian effects in female rats at the same LOAEL as liver effects in males was noted as a second reason to consider the rodent studies as an appropriate basis for deriving the pRfD.

Table 1. Summary of NOAELs, LOAELs, BMDLs, and Critical Effects for Key and Supporting C8 Studies								
	Species	Sex	NOAEL	LOAEL	BMDL	Critical Effect		
Key Studies								
Palazzolo et al. (1993)	Rat	M	0.47	1.44	1.3	Liver		
York et al. (2002)	Rat	M	None	1	0.42	Liver		
Riker Laboratories	Rat	F	None	1.6	1.6	Ovary		
(1983)		M	1.3	14	0.73	Liver		
Thomford et al. (2001)	Monkey	M	None	3-10	None	Liver		
Supporting Studies								
Goldenthal et al. (1987a)	Rat	M	0.56	1.72	0.44	Liver		
Goldenthal et al. (1987b)	Monkey	M,F	3	10	Not done	Clinical signs		

Some panelists favored choosing the monkey study as the critical study, due to the closer biological relationship with humans as opposed to rats. It was also noted that the observed increase in liver weight in monkeys may not be related to peroxisome proliferation and, therefore, may be more relevant for human health risk assessment. Other panelists disagreed, pointing to the uncertainties in dosing and effects, the small number of animals per dose group, and the unclear boundary between NOAEL and LOAEL values. Also, it was noted that the monkey study could not be considered the critical study because the 90-day, two-generation, and two-year rat studies all have LOAEL, NOAEL, and /or BMDLs below the LOAEL range identified in the monkey study, and therefore based on selection of the critical study with the lowest adequate NOAEL/LOAEL boundary would support the use of the rodent studies.

The panel considered whether it would be better to base the pRfD on a NOAEL or on a BMDL. Some panelists thought a NOAEL basis is a simpler concept and would be easier to explain to the public. Others responded that the BMDL captures more information from the entire study (e.g., reflects information from the full dose-response curve, and variability in the dose-response data) and therefore is the better choice as the basis for the quantitative dose-response assessment. Another panel member mentioned that a NOAEL is not a "no effect" level, rather it reflects the proportion of the responding population that can physically be observed in an experimental situation. Therefore, the size of the population is important. The panel agreed to not rule out using either a NOAEL or BMDL, but instead to focus on the quality of each study and the lowest critical effect level it provided.

The panel noted the unusually good agreement of the NOAELs and LOAELs from all the studies. The lowest NOAEL observed in one of the potential key studies was 0.47 mg/kg-day, from the 90-day rat study by Palazzolo et al. (1993). The lowest LOAEL observed in a key study was 1 mg/kg-day from the rat two-generation study (York et al., 2002). This study did not test doses low enough to identify a NOAEL; however, the BMDL value estimated for this study, 0.42 mg/kg-day, was essentially the same as the observed NOAEL from the 90-day study. Therefore, the panel agreed that the BMDL was an appropriate NOAEL surrogate for the two-generation study. The ovarian stromal hyperplasia reported in the chronic rat study (Riker Laboratories, 1983), provided a higher LOAEL than the two-generation study, and the BMDL for this effect resulted in the same value as the LOAEL. This demonstrates that the liver endpoint is the critical effect, because it occurs at lower doses.

<u>Outcome</u>: Because of the consistency in NOAELs/LOAELs and critical effect in all the key studies, the panel concluded that all studies could be considered co-critical studies and that all provide important information for human risk assessment. However, the panel unanimously agreed that the NOAEL surrogate from the two-generation study, a BMDL of 0.42 mg/kg-day, should serve as the point-of-departure for the pRfD. This value was selected since it represented the lowest NOAEL or BMDL, and provided the added consideration of having evaluated reproductive and developmental effects.

Uncertainty Factors

If adequate human data are available, these data are used as the basis for noncancer risk factor development. Otherwise, animal study data are used, along with a series of professional judgments that are incorporated into the risk factor as "Uncertainty Factors" and account for an assessment of the relevance and scientific quality of the experimental studies. There are five different uncertainty factors commonly used to address issues of biological variability and uncertainty. Two factors (Interspecies and Intraspecies) are used to address variability or heterogeneity that exists between animals and humans, and within different human populations. Three factors (Subchronic, LOAEL, Database) are used to address lack of information. Typically, the maximum total uncertainty factor that EPA will apply is 3000. If all five areas of uncertainty/variability are present warranting a total UF of 10,000, then EPA generally concludes that the uncertainty is too great to develop an RfD. The panel discussed each area of variability or uncertainty separately. A short introduction to each area of uncertainty is provided below to aid the reader in evaluating the discussions of the panel.

Intraspecies Variability (UF_H): This factor accounts for the natural differences that occur between human subpopulations and for the fact that some individuals may be more sensitive than the average population. This factor is composed of two subfactors – one to account for toxicokinetic differences (how the body distributes and metabolizes the chemical) and one to account for toxicodynamic differences (how the body responds to the chemical). If no information is available on human variability, then a default value of 10 is used. If adequate information is available on one of the two subcomponents, then this information is used along with a default value of 3 for the remaining subfactor. If data are available to adequately describe human variability in both subfactors, then actual data may be used to replace default values. In addition, if a RfD is based on human data gathered in the known sensitive subpopulation, a value of 1 may be chosen for this factor.

The panel discussed the lack of available data describing human variability. One panelist suggested a comparison of human C8 blood levels and values from the animal studies. The highest human serum C8 level reported was 111 ppm, but the average was approximately 5 ppm. No effects were noted in the human subject with the highest blood level. Thus, at least some people achieved serum C8 levels equivalent to those that resulted in adverse effects in animal studies.

As noted in the discussion of the human data above, the panel acknowledged gaps in the data on human variability and inability to define the most sensitive subpopulation, and therefore concluded that the default value of 10 was appropriate for this factor.

<u>Interspecies Variability (UF_A):</u> This factor accounts for the differences that occur between animals and humans and is also thought to be composed of subfactors for toxicokinetics and toxicodynamics. If no information is available on the quantitative differences between animals and humans, then a default value of 10 is used. If information is available on one of the two subcomponents, then this information is used along with a default value of 3 for the remaining subfactor. If data are available to adequately describe variability in both subfactors, then actual data may be used to replace default values. In addition, if a RfD is based on human data, then a value of 1 is appropriate for this factor.

One panelist mentioned that EPA has often used a UF_A value of 3 in other assessments when extrapolating monkey data to humans, because the kinetics and dynamics of monkeys are assumed to be similar to humans. This assumption is based on the fact that rhesus monkeys and macaques share a 92% genetic homology with humans and because monkey studies are able to detect a much broader range of clinical findings and more specific histopathology than rodents. In addition, studies on other chemicals in which a good database exists in rodents, monkeys and humans demonstrate that results in monkey studies parallel the human effects more closely than results in rodent studies.

Another panelist agreed and said the half-life of chemicals in monkeys was usually closer to humans than to rats. Other panelists responded that for C8, the half-life in monkeys is about 30 days; and this is much less than the C8 half-life in humans, which is estimated to be greater than one year. It was noted, however, that data on C8 half-life in humans is limited.

Because no data are available to warrant moving from the default, the panel unanimously agreed that a UF_A value of 10 is appropriate with either the rat or monkey toxicology studies.

Subchronic to Chronic Extrapolation (UF_S): Because the RfD protects for a lifetime exposure, this factor is applied when the database lacks information on the health effects of the chemical following a chronic exposure. Two issues are considered when making judgment on the use of this factor – are there data demonstrating that different health effects are expected following chronic exposure than subchronic exposure, and are there data demonstrating that the observed health effects progress in severity as exposure duration increases? If the database contains no information on chronic exposure, a default value of 10 is often applied, unless other data suggest a lack of progression with exposure duration. If the database contains adequate chronic bioassays, then a value of 1 is appropriate. If there are data addressing only one of the two issues, then a default of 3 may be applied.

It was noted that the database for C8 contains an adequate chronic rat study (Riker Laboratories, 1983). In addition, a second chronic study (Biegel et al., 2001) was available, although this study focused primarily on tumorigenic mechanisms in rats. In addition, for the purpose of evaluating uncertainty factors, the human occupational studies were considered by the panel to be informative on the response (or lack thereof) of humans following long-term exposure. The database demonstrates that liver

toxicity was the more sensitive endpoint in both subchronic and chronic studies. In addition, the database clearly demonstrates that liver toxicity does not progress in severity following chronic exposure. This conclusion is supported by the observation that the subchronic studies identified lower NOAELs for liver toxicity than the chronic studies. One panelist noted that the liver effect in rat progresses to cancer. However the panel concluded that the cancer effect was due to the peroxisome proliferation mechanism (as discussed below in the discussion of the cancer risk assessment). Based on these considerations, the panel unanimously agreed that a UF_S value of 1 is appropriate for the rat studies.

The panel also discussed whether a different value for UF_S would be appropriate if the monkey study had been used as the critical or co-critical study. One panelist observed that there were no data in monkeys regarding the progression beyond 26 weeks; another responded that there was no reason to think the effects in monkeys would be any more progressive than those in rats. Another panelist suggested that the toxicity of C8 in humans does not appear to be progressive. However, the panel agreed that there was some inherent uncertainty in the monkey study to justify use of the value of 3 for UF_S if the monkey study were the critical study.

LOAEL to NOAEL Extrapolation (UF_L): Because the RfD is considered to be a subthreshold value that protects against any adverse health effects, this factor is applied when the database lacks information to identify a NOAEL. If the database does not identify a NOAEL, then a default of 10 is used for this factor. If a NOAEL is used, a value of 1 is appropriate. Often, if the database does not identify a NOAEL, but the adverse effects observed are of minimal severity, then a default of 3 will be considered appropriate for use of a "minimal LOAEL".

Several of the studies considered as co-critical identified NOAELs; the lowest NOAEL is 0.47 mg/kg-day from the 90-day study. Also, the BMDL estimated for the two-generation study was essentially the same as the observed NOAEL from the 90-day study. These NOAELs and BMDLs were based on well-conducted studies and their use as a basis of the pRfD is consistent with standard practice. Therefore, the panel had confidence that the C8 database has identified the threshold for toxicity in rats, and it unanimously agreed a UF_L value of 1 is appropriate for the critical effect in the rat studies.

The panel also considered the value of UF_L that would be appropriate if the monkey study were to be used as the critical study. Because there is no clear NOAEL value, the panel agreed that a value of 1 was not appropriate. However, because the effects seen at the low dose were limited to mild increases in liver weight without accompanying changes in histopathology, or any other effect, the low dose was considered to be a minimal LOAEL. Therefore, the panel agreed that a UF_L of 3 would be appropriate if the monkey study were to be used as the critical study.

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¹ EPA is currently discussing the application of UF_L when using a BMDL. A BMDL value represents the lower limit on the dose that should cause 10% of the experimental animals to respond with the effect that is being modeled. Because animal studies typically cannot detect a response less than 10%, an experimentally derived NOAEL also represents the dose that causes 10% of the animals to respond. For this reason, EPA has historically considered a BMDL to be a NOAEL surrogate and selected a UF_L value of 1 when a BMDL is used. Although EPA does not have official guidance on this issue, recent discussions in the agency suggest that if the effect being modeled for the BMDL is adverse, then the BMDL should be considered as a LOAEL. Currently, BMDLs are being evaluated on a case-by-case basis, considering the nature of the effect being modeled and the relationship of the estimated BMDL to observed NOAELs.

<u>Database (UF_D)</u>: The database for deriving a high confidence RfD includes two chronic bioassays by the appropriate route of exposure in different species, one two-generation reproductive toxicity study, and two developmental toxicity studies in different species. The minimal database required for deriving a RfD is a single subchronic bioassay, that includes a full histopathology examination. The database factor is used to account for the fact that a potential health effect may not be identified if the database is missing a particular type of study. This factor may also be used if the existing data indicate the potential for a heath effect that is not fully characterized by the standard bioassays, for example neurotoxicity or immunotoxicity. If the database is complete, a value of 1 is appropriate. If only the minimal database is available, then a default of 10 is used. A value of 3 may be used if the database is missing one or two key studies.

The panel agreed that the oral database for C8 is complete. For the purpose of evaluating uncertainty factors, the panel felt that the human occupational studies provided sufficient information on the effects of long-term exposure in humans to function as a chronic bioassay. In addition, the consistency between the monkey and rat subchronic studies provides confidence that non-rodent species respond similarly to rats and that liver is a sensitive target organ in all species. Furthermore, a developmental toxicology study indicated that such effects only occurred at high concentrations, and reproductive effects were monitored in the 2-generation reproductive study.

Therefore, the panel unanimously concluded that a UF_D value of 1 is appropriate with either the rat or monkey toxicology studies selected as the critical study.

Outcome: The summary of the panel's unanimous conclusions regarding individual and composite uncertainty factors is presented in Table 2 below. The composite uncertainty factor is obtained by multiplying the individual factors. (Note, that following EPA convention, an uncertainty factor of 3 actually represents the log of the halfway point between 1 and 10. Therefore multiplying half-log values of 3 results in a full log value of 10, rather than 9 as would be expected for numeric multiplication.)

Table 2. Panel Recommendations of UF Selection for Oral pRfD						
Study	<u>UF</u> _H	<u>UF</u> _A	<u>UF</u> _L	<u>UF</u> _D	<u>UF</u> s	Composite UF
All Rat	10	10	1	1	1	100
Monkey	10	10	3	1	3	1000

Oral Reference Dose (RfD)

The final value of the RfD is obtained by dividing the point-of-departure by the composite uncertainty factor. As discussed above, the point-of-departure selected by the panel is the BMDL of 0.42~mg/kg-day estimated from the rat two-generation study (York et al., 2002) and the composite factor is 100. Therefore, the resulting pRfD is $0.42 \div 100$, or 0.0042~mg/kg-day. Because of the lack of precision inherent in the RfD, only one significant figure is appropriate; therefore, this value is rounded to 0.004~mg/kg-day.

For comparison purposes, the panel considered the pRfD values that would result from choosing alternative NOAELs or BMDLs as the point of departure. This analysis is presented in Table 3 below:

Table 3. Comparison of pRfDs Derived Using Different Studies							
Study	<u>UF</u>	NOAEL	RfD	BMDL	<u>RfD</u>		
Palazzolo et al. (1993)	100	0.47	0.005	0.72	0.007		
Riker Laboratories (1983)	100	1.3	0.01	0.73	0.007		
York et al. (2002)	100			0.42	0.004		
Thomford et al. (2001)	1000	3-10 (LOAEL)	0.003-0.01				

Based on this review table developed by the panel, the pRfDs that could be derived from the C8 oral database range from 0.003 to 0.01 – at most a factor of 3 separates the different potential pRfDs. Considering that the definition of the RfD states that the RfD incorporates uncertainty spanning an order of magnitude (a 10-fold variation), the panel noted that close agreement of the potential pRfD values provides added confidence in the derived pRfD of 0.004 mg/kg-day.

Noncancer Assessment: Review of the Dermal Studies

(Note: Dr. Seed abstained from voting during this part of the meeting)

The data on C8 by the dermal route of exposure are limited. Other than acute lethality, skin sensitization, and irritation studies, the dermal database consists of only a single 2-week study.

Kennedy et al. 1985

This is a two-week study in male rats in which animals had C8 applied to the skin for 6 hours/day, 5 days/week at doses of 0, 4.2, 42, and 420 mg/kg-day. Although this is a short-term study, it is the only candidate for possible use in determining a reference dose for the dermal route of administration. The primary effects observed were increased liver weight and liver pathology. A panelist noted that the study design prevented animals from ingesting the dermally-applied material. Although the amount of material inhaled was considered to be low, some inhalation almost certainly occurred in the dosed animal because the control animals had detectable C8 blood levels. It was also noted that the consistency of the material applied to the animals varied among the dose groups, depending on the concentration of C8 in the material matrix. In all instances the amount of material on the skin was considerably thicker than a monolayer, and therefore, the applied doses might not reflect accurately the absorbed doses of C8 in this study.

Key Panel Discussion Points: One panelist stated that this study could provide potentially useful information because systemic effects are observed at dose levels below those which cause portal of entry effects (skin irritation). The panel discussed whether it would be appropriate to extrapolate the results of this study to longer durations in order to derive a dermal pRfD. The panel concluded that such extrapolation would not be advisable because of the possibility of unpredictable longer-term dermal effects. One panelist asked if route-to-route extrapolation could be done from the oral studies

to estimate a dermal NOAEL or LOAEL. Other panelists thought this would not be possible due to uncertainties in the C8 toxicokinetics by oral versus dermal exposure routes. For example, enterohepatic circulation is known to occur following oral exposure, but would not occur following dermal exposure. Therefore, the toxicokinetics of C8 is different between the two routes of exposure. Regardless of the route of entry, C8 is not metabolized. Furthermore, no data on the dermal absorption rate were identified. One panelist noted that if the findings from this study were used to determine a reference dose, the resulting value would be higher than the reference dose obtained from the oral studies. Therefore, using oral studies to set the reference dose would be adequately protective, of systemic exposure via the dermal route. Another panelist agreed, stating that no dermal reference dose should be identified at all, and that a specific reference dose for dermal exposure was not needed.

<u>Outcome</u>: The panel agreed unanimously that this study should not be used to determine a dermal pRfD because of uncertainties inherent in the study design as noted in the discussion.

Noncancer Assessment: Review of the Inhalation Studies

(Note: Dr. Seed was absent during this part of the meeting)

The data on C8 by the inhalation route of exposure are limited. Other than acute lethality studies, the inhalation database consists of a 2-week study and a developmental toxicity study.

Kennedy et al. 1986 and Staples et al. 1981

Two inhalation studies were discussed as potential candidates for deriving the pRfC. Kennedy et al. (1986) reported a two-week study in male rats in which animals were exposed head-only 6 hours/day, 5 days/week to C8 air concentrations of 0, 1, 7.6, or 84 mg/m³. The primary effects observed in this study at the mid-concentration included increased absolute and relative liver weight, supported by clinical chemistry and histopathology findings. The high concentration resulted in severe toxicity, including mortality in one rat. Other findings at the high concentration group were increased lung and testes weight. A concentration-dependent increase in the incidence of nasal and ocular discharge was noted.

A second potential critical study for deriving the pRfC was a developmental toxicity study by Staples et al. (1981). Pregnant rats were exposed whole-body 6 hours/day on gestation days 6 to 15 to C8 air concentrations of 0, 0.14, 1.2, 9.9, and 21.0 mg/m³.

The panel agreed the Kennedy two-week study provided the highest quality data for possible determination of critical effects and provided a slightly lower NOAEL/LOAEL boundary, even though both studies used similar air concentrations. In addition, the Kennedy et al. (1986) study evaluated a broader array of systemic endpoints, and included a histopathology examination.

In describing their initial review of the study, *TERA* noted that EPA's RfC methodology states that the air concentrations to which animals are exposed are to be converted to "Human Equivalent Concentrations (Conc_{HEC})" by applying dosimetric adjustments (USEPA, 1994). Dosimetric adjustments account for the different structure and surface area of animal respiratory tracts compared with humans. Different dosimetric adjustments are applied depending on where effects are observed. For example, a different dosimetric adjustment will be applied for liver effects than will be applied for lung effects. *TERA* noted that the key piece of data needed to calculate the Conc_{HEC} is a description of the particle size distribution (i.e., the mass median aerodynamic diameter and geometric standard

deviation or GSD). Data available from the published study did not provide complete information about the mass median aerodynamic diameter for the low-concentration group, or GSD for any exposure group. In order to facilitate the discussion of the study, *TERA* presented human equivalent concentrations for liver (extrarespiratory) and lung (pulmonary) effects from this study assuming either a monodisperse particle size distribution or a polydisperse particle size distribution. These results were presented to the panel as shown in Table 4 below.

Table 4. Preliminary Conc _{HEC} Calculations from Kennedy et al. (1986)							
Study Concentration ^a	GSD = 1.3 (Mo	GSD = 1.3 (Monodisperse) GSD = 3 (Polydisperse)					
	Liver Lung Liver Lung						
1.0	0.6	0.018	0.5	0.09			
7.6	4.6	0.14	4.0	0.70			
84 67.7 17.7 46.9 7.4							
a. All values are presented in units of mg/m ³ .							

<u>Key Panel Discussion Points</u>: It was noted that the inhalation database does not meet the minimum database requirements for determining an RfC of one subchronic 90-day study that includes histopathology of the respiratory tract, but that the consent order required a pRfC in order to set air screening levels. One panelist stated that it was not appropriate to extrapolate from oral studies to derive a RfC because of the absence of data on toxicokinetics differences between these routes (e.g., effects of enterohepatic circulation, or absorption).

One panel member indicated that the data needed to calculate the Conc_{HEC} (i.e., the mass median aerodynamic diameter [MMAD] and geometric standard deviation [GSD]), but not reported, in the published study could be made available to TERA after the meeting. The panel agreed that these data should be provided to TERA, for calculation of the appropriate Conc_{HEC} following the meeting. The panel then discussed whether the lung or the liver was the critical organ, recognizing that the final designation of critical effect could not be made until the correct Conc_{HEC} is calculated. TERA raised the question of whether the reported increases in the incidence of nasal and ocular discharge should be considered an adverse effect. It was noted that this effect is not uncommon for the exposure protocol that was used, and the effect was seen in all groups. It was further noted that C8 is not an irritant, and that no nasal histopathology was observed in exposed animals. In selecting critical study concentrations the panel discussed the lung effects at higher doses. One panel member suggested that at the high concentration the overt pulmonary toxicity was observed due to the large particle burden. Uncertainties in interpreting the lung effects were raised by the panel. One panelist noted that the studies were too short to determine what effect chronic exposure would have on the respiratory tract. Another suggested that existing human data associated with the human study reports discussed earlier (pulmonary function testing of workers, etc.) might be useful in determining NOAEL/LOAEL values. After this discussion, the panel considered the study concentration of 7.6 mg/m³ to be the NOAEL for pulmonary effects, with the LOAEL of 84 mg/m³.

The panel next discussed the liver effects. It was noted that the observed increases in liver weight were consistent with the effects observed in the oral studies. Another panel member noted the increased alkaline phosphatase (AP) values observed at the higher doses were not necessarily the result of the types of liver effects seen in the oral and dermal studies, since increased AP levels often reflect disorders of biliary flow. One panelist questioned the ability of the study to detect systemic effects given the short exposure period and the kinetics of the compound; however, another panelist replied that the half-life of C8 in rats is 5 to 7 days, and the study design would have allowed achievement of

steady-state concentrations in the blood. The panel considered the study concentration of 1.0 mg/m^3 as the NOAEL for liver effects. However, one reviewer suggested that if the liver effects are found to be the critical effect based on the Conc_{HEC}, then benchmark concentration modeling should be conducted before assigning a critical effect level.

The panel considered the appropriate uncertainty factors for a pRfC, noting that the final choice of an appropriate value for some areas of uncertainty may change depending on whether lung or liver effects are found to be critical. (Note to the reader: Essentially the same areas of uncertainty are considered in developing a RfC as for the RfD. For a full explanation of the purpose for each factor, see the earlier discussion.) For the same reasons as discussed for the pRfD, the panel unanimously agreed that a value of 10 was appropriate for UF_H. When considering interspecies extrapolation, it is generally considered that the dosimetric adjustments used to derive the Conc_{HEC} account for the toxicokinetic differences between animals and humans. Therefore, the uncertainty factor only needs to address the toxicodynamic differences. Since there are no data regarding dynamic differences between rats and humans, the panel agreed that the default value of 3 was appropriate for UF_A. Since the Kennedy study identified a NOAEL, the panel unanimously agreed that a value of 1 was appropriate for UF_L.

The panel considered that two of the factors, UF_S and UF_D , were related to the decision of whether lung or liver is the critical effect. If liver effects are determined to be the critical effect, then at least one panelist felt that UF_S , could be addressed with an uncertainty factor of 1 because the oral studies provided enough information to be confident that the liver effects would not progress in severity following a chronic inhalation exposure. However, other panel members stated that there were insufficient data to assess whether liver would continue to be the critical effect or to provide information on how the respiratory tract would respond following longer-term inhalation exposures, and that a value greater than 1 for UF_S was needed. For the UF_S and liver as the critical organ, the panel votes were 1, 3, or 10 with the majority choosing 3. If liver effects are determined to be the critical effect, then panelists were split on the value of the uncertainty factor for UF_D , choosing values of either 3 or 10 with the majority of the panel choosing 3. No unanimous consensus was reached on these two factors; however, a clear majority vote was reached on uncertainty factors of 3 each for UF_S and UF_D in reference to liver as the target organ.

If lung effects are determined to be critical, the panel was divided almost equally on the appropriate value for UF_S with opinions covering the full range of options from 1 to 3 to 10. Note however, that six scientists voted for a factor less than 10 (either 1 or 3) and five scientists voted for a value greater than 1 (3 or 10). Similarly, the panel was divided on the appropriate value for UF_D; panel opinions covered the full range of options from 1 to 3 to 10 with the majority of panelists choosing 3.

As noted above, after each discussion votes were taken on individual factors. These votes are shown in Table 5. Note that one scientist was reviewing the dosimetric adjustment calculations during this discussion and so was unable to vote on these UFs; also note that one more vote at any point in Table 5 would not have changed the final outcome. In addition, the panel did not reach consensus on the confidence in the RfC, with opinions ranging from "none" to "high" with the average being mediumto-low.

Table 5. Tally of Panel Votes for UF _S and UF _D							
		$\mathrm{UF_{S}}$			UF _D		
Factor	1	3	10	1	3	10	
Liver as critical	1	6	1	0	6	2	
Lung as critical	3	3	2	1	5	2	

Outcome: One panelist reminded the group that the purpose of Kennedy et al., (1986) was to identify the inhalation hazard, not to look closely at NOAEL, LOAEL, etc. A prospective inhalation study designed to look more closely at the NOAEL/LOAEL aspects, to evaluate lesions as a function of exposure time, and to evaluate tissues of the respiratory tract using up-to-date methodology would be valuable and would allow a more focused evaluation of the RfC. Nonetheless, the panel agreed that a pRfC could be developed, but this agreement was not unanimous. The panel also recommended that *TERA* obtain additional data on the particle size GSD value to determine the Conc_{HEC} corresponding to the NOAEL before determining whether the pulmonary or the hepatic effects are considered critical. If the liver effects are determined to be the critical effect, then BMD modeling should be done. The composite uncertainty factor was expressed as a range of 30 to 3,000. The final pRfC is presented in the Post Meeting Action Items.

Cancer Assessment

(Note: Dr. Seed abstained from voting during this part of the meeting)

U.S. EPA's 1999 Guidelines for Carcinogen Risk Assessment were used to frame the discussion of C8 carcinogenic potential. *TERA* opened the discussion with a short introduction to these guidelines, highlighting the recent focus on evaluation of the mode of action data in developing a weight of evidence characterization, and in deciding the most appropriate dose-response approach, linear or margin of exposure (MOE). It was noted that the EPA's 1999 guidelines would be used as the basis for the deliberations of the panel.

Cancer Hazard Identification and Mode of Action

The panel discussed the evidence for C8 carcinogenicity in humans and agreed that the human carcinogenicity evidence is inconclusive. Although four prostate tumors were reported in retired workers, three of these four cases now are known to have had minimal or no C8 exposure. (See Human Studies section for more detailed discussion.)

The panel noted that two animal carcinogenicity studies had been conducted. The first study (Riker Laboratories, 1983) reported treatment-related increases in Leydig cell adenomas and mammary gland fibroadenomas. The second study (Biegel et al., 2001) reported treatment-related increases in tumors in the liver, Leydig cells, and pancreas. Panelists noted that the tumors identified in the Biegel et al. (2001) study correspond to the triad of tumors associated with some chemicals that cause peroxisome proliferation. Other panelists agreed and suggested that a further examination of the data may indicate that this triad of tumors can be best addressed using a MOE approach. The panel also noted that the mammary fibroadenomas may require the default linear model because, following U.S. EPA cancer guidelines, no actual mode of action data for C8 and this tumor type are available to warrant moving from the default assumption. Each of the four types of tumors found in the two C8 animal

carcinogenicity studies was then discussed in detail with regard to the weight of the evidence for the mode of action, and the evidence supporting a linear or MOE dose-response assessment approach. Listed below are the outcomes and discussions for each tumor type.

Liver tumors

Key Panel Discussion Points: The discussion on liver tumors focused on the role of peroxisome proliferation as the mode of action for the observed liver tumors. In relating this liver tumor effect to humans, one panelist said humans are much less sensitive to peroxisome proliferation than rats. Another panelist noted that IARC's approach for clofibrate and other non-genotoxic peroxisome proliferation chemicals was to assume that the mode of action was not relevant to humans if no evidence of peroxisome proliferation was observed in humans. Another panelist said that although rats may be more sensitive than humans from a toxicodynamic standpoint (due to interspecies differences in receptors), humans may be more sensitive from a toxicokinetic standpoint, since they clear C8 more slowly than rats. As a result, the panel member suggested that these two considerations would tend to decrease overall differences in species sensitivity. On the other hand, a panel member noted that no increased incidence of tumors have been found in people taking clofibrate, a known peroxisome proliferator, which suggests that humans are much less sensitive to peroxisome proliferation than rats and they may have no response at all. Based on these data, the panel member suggested that the lack of tumor development in humans exposed to C8 should not be discounted. The panel discussed differences in results between the two cancer studies. One panelist noted the studies have differences in their internal delivered doses because of differences in the animal diets. This could explain the difference noted in toxic effects.

<u>Outcome</u>: The majority of the panel agreed that the data indicate peroxisome proliferation is the mode of action for the liver tumors, and that although the liver tumor response is not likely to be quantitatively similar between rats and humans, the use of the liver tumor response data for human health risk assessment cannot be totally discounted. However, other scientists indicated that based on the lack of peroxisome proliferation in the non-human primate studies, the rodent liver tumors are not relevant at all to humans.

Leydig Cell Tumors

<u>Key Panel Discussion Points</u>: In reviewing the summary tables prepared for the meeting, one panelist noted that Leydig cell hyperplasia should be evaluated. In response, the hyperplasia data from Biegel et al. (2001) was reviewed by the panel. The panel developed Table 6 to facilitate the comparison on hyperplasia and tumorigenic outcomes.

Table 6. Summary of Beigel et al., 2001 Leydig Cell Data						
	Pair fed controls	300 ppm				
Liver carcinomas/adenomas	3/79	10/76*				
Leydig ademonas	2/78	8/76*				
Pancreatic carcinomas/adenomas	1/79	8/76*				
Leydig cell hyperplasia	26/78	35/76				

The panel noted that no significant increase in Leydig cell hyperplasia was apparent from these data; however, due to different survival times between the two groups (C8 treated animals survived longer) a false positive effect could have occurred because older animals would have more time to develop naturally occurring tumors. It was noted that a more formal analysis would be needed to determine whether the incidence of Leydig cell tumors would still be increased after adjusting for differences in survival, but the formal statistical analysis was too complex to complete during the meeting.

The panel discussed the role of peroxisome proliferation as the mode of action of Leydig cell tumors. Specifically, the panel discussed a workshop publication (Clegg et al. 1997) that evaluated the seven known modes of action for Leydig cell tumors. Most of the modes of action involve altered hormonal response in response to peroxisome proliferation, including increased estradiol via hepatic aromatase and binding to the TGF α receptor or elevations in leutinizing hormone to compensate for the testes becoming less responsive to this hormone. One panelist emphasized that the monkey study (Thomford et al., 2001) showed no effects in the testes, even though the animals were dosed at C8 levels high enough to cause major weight loss and mortality. This panelist suggested that this indicates the Leydig cell effects seen in rats are unlikely to occur in primates. This panel member also noted that no increased estradiol was noted in the monkeys.

One panelist observed that Leydig cell tumors were a classic response to peroxisome proliferation but the available studies do not provide positive evidence, such as increased estradiol levels, that peroxisome proliferation is the operative mode of action. The panelists agreed that while data gaps exist, a peroxisome proliferation mode of action was a reasonable assumption. One panelist stated that whatever the MOA was, it was not genotoxicity.

The panel agreed unanimously that for Leydig cell tumors:

- All 7 possible mechanisms for Leydig cell tumors are non-linear; therefore a non-linear dose-response approach is reasonable;
- Humans have a low incidence of these tumors;
- The monkey study did not demonstrate Leydig cell pathology or increased estradiol;
- Leydig cell tumors are a known tumor type for other peroxisome proliferators;
- Humans do not develop Leydig cell tumors following exposure to other known peroxisome proliferators such as clofibrate;
- Regardless of the actual mode of action, it is likely to be non-genotoxic.

<u>Outcome</u>: The panel agreed that based on the absence of genotoxicity, the Leydig cell tumors were likely to be caused by a non-genotoxic mechanism. The panel further agreed that if sufficient evidence were available to show increased estradiol levels (i.e., secondary to peroxisome proliferation) as the mechanism for the observed tumors, then the mechanism would be non-genotoxic and would not be quantitatively similar or possibly not relevant at all to humans. However, without this evidence this effect can not be totally discounted.

Pancreatic tumors

<u>Key Panel Discussion Points:</u> Since the tumor results from the Beigel et al., (2001) were not provided in the summary table distributed to the panel prior to the meeting, the pancreatic tumor data from this study were presented as a table at the meeting (see Table 7 below):

Table 7 Biegel Study: Pancreas Tumors						
Control pair-fed control 300 ppm						
Hyperplasia	14/80 (18%)	8/79 (10%)	30/48* (40%)			
Adenomas	0/80	1/79	7/76*			
Carcinomas	0/80	0/79	1/76			

One panelist described an analysis that had been done to compare the two cancer studies with regard to the pancreatic tumors. This panelist noted that although the first study (Riker Laboratories, 1983) did not report pancreatic tumors or hyperplasia, the second study (Biegel et al., 2001) did. However, this panel member also noted that the studies were not inconsistent because of the different definitions of adenoma versus hyperplasia based on pancreatic cell size used by the respective investigators. Also, the criteria for separating hyperplasia from adenomas is based on lesion size. Both studies were qualitatively similar with a number of larger lesions (adenomas) found in the Biegel study. Another scientist commented, when the two studies were recently compared by a group of pathologists using current criteria, there was a consistency in a pancreatic response; however, there was not an increased number of adenomas found in the earlier study. Instead, an increase in hyperplastic nodules of the acinar pancreas was found, which is consistent with the Beigel study. However, even though the dietary dose was the same (300 ppm), the Riker Laboratories study rats did not develop these hyperplasias into adenomas to the extent that occurred in the Beigel study.

With regard to the potential mode of action, one panelist suggested that the persistent increase seen in cholecystokinin and increased bile acids may be involved in the MOA, but the evidence in rats, monkeys and humans does not support this hypothesis. When a panelist asked if a strong case could be made that the pancreatic tumors resulted from peroxisome proliferation, several panelists responded no. Another added that while some peroxisome proliferation agents cause the triad of tumors seen with C8, not all do. Another panelist added that no pancreatic, liver, or testes hyperplasia was noted in monkeys at the time of sacrifice.

<u>Outcome</u>: The panel agreed that the evidence was not sufficient to demonstrate the MOA for pancreatic tumors, but enhanced cell proliferation (hyperplasia) was likely to be involved. The MOA appears to be non-genotoxic based on the results of genotoxicity bioassays.

Mammary Fibroadenomas

<u>Key Panel Discussion Points:</u> The panel considered whether the fibroadenomas observed in the Riker Laboratories study were a real treatment-related effect, or an artifact of classification, since other mammary tumor types observed in this study showed no clear relationship with dose. Table 8 below shows the data for several types of mammary tumors from this study:

Table 8. Riker Study: Mammary Tumors					
	Control	30 PPM	300 PPM		
Adenomas	7%	0	0		
Adenocarcinomas	15%	31%	11%		
Carcinomas	2%	0	0		
Fibroadenomas	22%	42%	48%*		

One panelist suggested that even though fibroadenomas were statistically significant, when all mammary tumor types are combined, they are not likely to be significant. It was noted by the panel that the individual incidence data from the study would need to be examined to determine the combined incidence of all mammary tumor types, rather than adding the percentages from each category. The panel discussed the histological basis for reporting separately fibroadenomas versus other types of mammary adenomas. A panelist suggested that since fibroadenomas do not progress to the other types it is correct to report them separately. Another said that the National Toxicology Program (NTP) reports fibroadenomas combined with adenomas.

The panel also discussed potential modes of action for mammary tumors. Increased estradiol was proposed as a possible MOA for the induction of hyperplasia and tumor formation, but the panel did not believe the data were sufficient to demonstrate this proposed mode of action. A panelist asked if a linear assessment could be done to help decide the importance of the effect. Another responded that the data were not adequately fit by any of the acceptable dose-response models, so a quantitative dose-response assessment was not reported for this data set.

<u>Outcome</u>: The panel agreed the data are not adequate to demonstrate a MOA; however based on the negative genotoxicity assays, C8 is unlikely to be genotoxic. Several panelists were not convinced the data demonstrated any real tumorigenic effect.

Cancer Dose-Response Assessment

After evaluating the relevance of each tumor type to humans, and the potential mode of action, the panel members were asked to recommend a dose-response approach for each tumor type. In all cases the panel agreed unanimously unless noted otherwise. For the liver tumors, the panel agreed that the MOE approach was most appropriate. For the remaining tumor types, the panel agreed that both linear and MOE approaches were appropriate, since the mode of action was not considered to have been adequately demonstrated for any of these three tumor types. All panel members agreed with these conclusions, except for the Leydig cell tumors, where one panel member argued that only an MOE approach should be used.

For the liver tumors, the MOE approach was selected. Since the MOE analysis often uses the benchmark response for a precursor as the basis of deriving a point of departure, the panel judged the pRfD for liver effects as sufficiently protective of potential liver carcinogenicity.

For Leydig cell tumors, benchmark dose modeling was conducted to identify a point of departure for the linear and MOE assessments. The Point of Departure (POD) for Leydig cell was chosen by the panel from the BMD modeling output. The BMDL of 0.32 mg/kg-day was selected as the most appropriate basis for deriving the assessment.

The panel discussed the appropriate factors to apply to the BMR for completing the MOE assessment. The panel noted that EPA's 1999 guidelines have only recently begun to be applied, and that formal guidance or examples of the interpretation and default values to use in deriving the MOE are lacking. In discussing the important considerations for the MOE, the panel decided that the critical factors to be considered were for "Nature of Effect", Intrahuman sensitivity" and "Animal to Human Extrapolation". A summary of the factors chosen is shown in Table 9.

For the Leydig cell tumors, a factor of 3 for nature of effect was selected as the most appropriate value, since the observed effect was for benign tumors. A factor of 10 was selected for Intrahuman sensitivity. A factor of 3 was used for Animal to Human Extrapolation, since dosimetric adjustments were applied to the dose data used for the BMD modeling. This composite factor of 100 was further supported since these types of tumors, although common in rats, are found rarely in people. In addition, the mode of action is likely via peroxisome proliferation which is quantitatively much less important in humans. The panel agreed that the composite MOE of 100 was appropriate.

For the linear dose-response assessment for Leydig cell tumors the BMDL of 0.32 mg/kg-day was used to calculate an oral cancer slope factor as follows:

Slope factor = risk/dose = 0.1/0.32 = 0.31 per mg/kg-day

(Note: risk is numerically expressed as 0.1 because the BMDL is the point that represents a 10% increased in tumor incidence in accordance with EPA guidance.) BMD modeling failed for the tumor data for pancreatic tumors and mammary gland fibroadenomas. Therefore, the panel determined that the data for these two tumor types were not adequate to conduct a quantitative dose-response assessment.

Table	9.
Factors Used to Describe	Various Areas in the
Development of MOEs for	or Cancer Endpoints.

		Nature	Intra	Animal	Steepness	Total	
<u>Tumor</u>	Model	Of Effect	<u>Human</u>	to Human	of Slope	Exposure	<u>MOE</u>
Liver	MOE	1	10	10	NR	NR	100
Leydig	both	3	10	3	NR	NR	100
Pancreas	both	NA (can	not be mod	leled)			
Mammary	both	NA (can	not be mod	leled)			

NR = Not Relevant based on panel judgment; NA = Not Applicable

The panel also voted on confidence ratings for the cancer assessment. *TERA* noted that according to EPA guidance "high confidence" suggests that the assessment is unlikely to change with the availability of new data, while "low confidence" indicates that the assessment is likely to change with new data. Based on these criteria the panel voted on their confidence in the cancer assessment using either the pRfD for liver toxicity to adequately account for the liver cancer risk or using the assessment based on Leydig cell tumors. The panel voted as follows:

```
Liver pRfD = high (7 votes); medium-high (2 votes)
Leydig tumors = low (7 votes); low-medium (2 votes)
```

Therefore, the panel agreed that the oral pRfD for liver toxicity would be the basis for determining water and soil screening levels (which are based primarily on oral exposure) for the following reasons:

- high confidence in the pRfD (i.e., not likely to change in the future due to additional data collection);
- the pRfD would be protective against the quantitatively less sensitive and questionable relevance peroxisome proliferation-related liver cancer in humans;
- low confidence in the Leydig tumor analysis and questionable relevance to humans;
- limitations in study design, data quality, and data interpretation rendered difficult the determination of whether the reported increased incidence of pancreatic tumors or mammary tumors were related to C8 treatment, and did not allow the modeling of a point of departure that could serve as the quantitative basis for risk value development.

Screening Levels

(Note: Dr. Seed was absent during this part of the meeting)

The consent order required that screening levels be developed for drinking water, soil, and air. The panel followed the guidance provided by U.S. EPA's "Risk Assessment Guidance for Superfund" as further explained by both Region 3 and Region 9 risk-based concentration guidance. In cases where a conflict occurred between the guidance documents, Region 9 guidance was followed because it is more conservative, i.e. more health protective. For drinking water and soil, only ingestion and dermal absorption were considered as routes of exposure. EPA guidance indicates volatilization from water or soil should only be evaluated for chemicals with Henry's law constants greater than 10⁻⁵ and molecular weights less than 200. Since C8's Henry's Law constant is 10⁻¹¹ and its molecular weight is 431, volatilization was not evaluated.

As discussed above, the panel concluded that since both liver and Leydig cell tumors were potentially formed via nonlinear modes of action, and further since greater confidence was placed in the quantitative assessment based on the liver endpoint, the pRfD and pRfC for liver toxicity would be protective of potential cancer effects of C8. The panel considered that the linear extrapolation for Leydig cell tumors was too uncertain to be used with confidence and that the MOE approach based on the Leydig cell tumors gave essentially the same numerical value as that for the liver endpoint, but with less confidence. Thus, the pRfD and pRfC for liver toxicity, and "noncancer" equations were used for calculating screening levels. Screening levels are calculated following the premise that if lifetime exposure is equal to or less than the pRfD or pRfC, then no risk of deleterious effects is expected. Mathematically, this concept can be expressed by the following standard equation; the ratio of the measured or estimated exposure to the RfD is called the Hazard Quotient.

If Exposure \div RfD = 1 or less, then no risk of deleterious effects is presumed.

Using this concept, it is possible to estimate the concentration in media that results in a lifetime exposure equal to the pRfD or pRfC. These equations, from EPA Region 9's guidance on deriving risk based concentrations, are listed below:

Air Screening Level: [] $ug/m^3 = \frac{THQ \times RfDi \times BW \times AT \times 1000}{EF \times ED \times air IR}$

Note: RfDi (mg/kg-day) = RfC x $\frac{20\text{m}^3/\text{d} (IR)}{70 \text{ kg (BW)}}$

Soil Screening Level: $[] mg/kg = \frac{THQ \times AT \times BW}{EF \times ED \times [soil IR / RfD \times 10^{-6} + SA \times AF \times ABS / RfD \times 10^{-6}]}$

Water Screening Level: [] $ug/L = \frac{THQ \times AT \times BW \times 1000}{EF \times ED \times [water IR / RfD]}$

Where:

THQ = Target Hazard Quotient, assumed to be 1

RfDi = The RfC expressed in terms of dose, mg/kg-day

RfD = The oral reference dose estimated by the panel, 0.004 mg/kg-day

RfC = The inhalation reference concentration estimated by the panel, see below BW = Body weight, assumed to be 70 kg for adults and 15 kg for children AT = Averaging time, 10950 days, the exposure duration expressed in days

EF = Exposure Frequency, 350 days/year, the average number of days each

year people are exposed

ED = Exposure duration, 30 years, the average number of years people are

exposed

IR = Inhalation rate for air screening levels, 20 m³/day; Ingestion rate for soil

and.

Water screening levels, 200 mg/day soil ingested based on child exposure

and.

2 L/day water ingested based on adult exposure

SA = Surface area of exposed skin, $2800 \text{ cm}^2/\text{day}$

AF = Adherence factor, 0.2 mg/cm², the amount of soil that adheres to skin ABS = Skin absorption factor, specific factor not available for C8, assumed to be

0.1 for semi-volatile chemical per EPA guidance

The panel unanimously agreed that the equations, assumptions, and default exposure parameters described above were the appropriate choices for calculating screening levels for air, soil, and water. The following values are the screening levels estimated by the equations.

For air: 0.1-6.0 micrograms per cubic meter of air ($\mu g/m^3$) ambient air. Note that the panel considered this range to be interim until the additional work discussed for the RfC is completed. This range incorporates the range of possible NOAEL_{HEC}s estimated by *TERA* prior to the meeting as well as the range of composite uncertainty factors recommended by the panel. The final pRfC is discussed in the following section Post Meeting Action Items.

For soil: 244 miligrams per kilogram of soil (mg/kg) residential soil, rounded to 240 mg/kg.

For water: 146 micrograms per liter of water (μ g/L), rounded to 150 μ g/L.

2.3 POST MEETING ACTION ITEMS

The following activities were conducted after the CATT Toxicologists meeting.

Derivation of the pRfC for C8

The CATT panel could not develop a final recommendation on the pRfC or the air screening level during the May 6 and May 7, 2002 meeting. This was due to a lack of data necessary for these calculations. At the meeting, the panel chose the key study for risk factor derivation as the 2-week inhalation study by Kennedy et al. (1986) and voted upon the uncertainty factors. They directed the author, panel member Kennedy (DuPont), to (1) retrieve the standard deviation data for the absolute and relative liver weight data sets; and (2) to measure the particle size distribution in the exposure chamber and determine the corresponding standard deviation; and (3) to provide these data to DEP and to *TERA*. The panel directed *TERA* to utilize these data to develop the pRfC based on the most sensitive organ (liver or lung) and the air screening level based on USEPA Region 9 standard formulas.

During the meeting, the CATT panel agreed that the Kennedy et al. (1986) study was the most appropriate basis for deriving the pRfC, with the developmental study by Staples et al. (1981) providing support for the selected critical effect levels. The CATT panel identified a NOAEL for increased liver weight at the lowest study concentration of 1.0 mg/m³, with a LOAEL of 7.6 mg/m³. The NOAEL for lung effects was identified by the CATT panel as 7.6 mg/m³, with a LOAEL was 84 mg/m³.

In order to derive an pRfC, the reported study concentrations were converted to human equivalent concentrations (Conc_{HEC}), according to current U.S. EPA RfC methodology (USEPA, 1994). The calculation of the Conc_{HEC} requires two steps. First, the study concentration is adjusted from the exposure duration used in the experiment to an equivalent continuous exposure concentration (Conc_{ADJ}). Animals in this study were dosed for 6 hours per day, for five days, then not dosed for two days, and dosed again for five days and sacrificed at the end of the 12th day; hence, continuous exposure duration adjustment was made as follows:

Study concentration x (6 hours/24 hours) x (10 days/12 days) = $Conc_{ADJ}$

Second, the duration-adjusted concentrations (Conc_{ADJ}) were converted to human equivalent concentrations (Conc_{HEC}) to account for differences in the respiratory tract anatomy and physiology for the test species versus humans. This conversion is made as follows:

 $Conc_{ADJ} \times RDDR = Conc_{HEC}$

The RDDR is the Regional Dose Deposition Ratio calculated using U.S. EPA's RDDR software program (USEPA, 1994). The RDDR depends on the characteristics of the particle size distribution (e.g., mass median aerodynamic diameter, and geometric standard deviation), the test species and body weight, and the region of the respiratory tract (or extrarespiratory tissue target if applicable) affected by exposure. Appropriate particle size characteristics to use as inputs into the RDDR software were obtained from a recent communication from DuPont (see attached). For the Kennedy et al. (1986) study, the test sex and species was male rats. Since body weight data were provided in the study, these data were used directly in the RDDR program. The mean body weight data on day 5 of exposure was used for this calculation, rather than the study-day 10 body weight data. The day 5 body weights were

used because there was evidence of changes in body weight over the 12-day study period, and therefore, this value was judged as the best estimate of the mean body weight over the period of exposure.

The CATT panel considered two potential critical effects for deriving the pRfC; increased liver weight and overt toxicity secondary to pulmonary toxicity. The RDDR for extrarespiratory tissues was the most appropriate value to use in calculating human equivalent concentrations for assessing the liver effects. The RDDR program calculates values for a variety of different regions of the respiratory tract. The CATT panel agreed that the overt toxicity of C8 was likely due to particle overload, as supported by pulmonary edema in the acute study reported in the same paper (Kennedy et al., 1986). Therefore, the RDDR for the pulmonary region was selected as most appropriate respiratory tract region for calculating the human equivalent concentrations. The calculation of the human equivalent concentrations used in the dose-response assessment is summarized in Table 10.

Table 10. Calculation of Human Equivalent Concentrations for Kennedy et al. (1986)							
		Extrarespiratory		Pulmonary			
Study Concentration ^a	Conc _{ADJ}	RDDR ^b	Conc _{HEC}	RDDR	Conc _{HEC}		
1.0	0.21	2.956	0.62	0.513	0.11		
7.6	1.6	2.954	4.7	0.512	0.81		
84	17	2.973	52	0.521	9.1		
a. All concentrations reported in the table are in units of mg/m ³ .							
b. The RDDR values a	are taken from	the EPA RI	DDR Program	Output provided	in the		
attachment							

Benchmark Concentration Modeling

The CATT panel further recommended that benchmark concentration (BMC) modeling be performed for the increased liver weight endpoint from the Kennedy et al. (1986) study. The published version of the study did not provide standard deviations to accompany the group mean data, and therefore, BMC modeling could not be performed at the time of the CATT panel meeting. Subsequent to the meeting, the individual liver weight data for this study were obtained from DuPont (see attached). The individual animal data were used to calculate group mean and standard deviations. These data were then employed for the BMC analyses.

The modeling was conducted according to draft EPA guidelines (U.S. EPA, 2000) using Benchmark Dose Software (BMDS version 1.3.1), available from the U.S. EPA website (U.S. EPA, 2002). The endpoints of interest with respect to C8 liver toxicity were continuous rather than quantal (e.g., incidence data) in nature. Therefore the absolute and relative liver weight data sets were modeled using the linear, Hill, power, and polynomial models. An acceptable fit to the data was defined as a goodness-of-fit p-value greater than or equal to 0.1, or a perfect fit when there were no degrees of freedom for a formal statistical test of fit. Choice of 0.1 is consistent with current U.S. EPA guidance for BMD modeling (U.S. EPA, 2000). Goodness-of-fit statistics are not designed to compare different models, particularly if the different models have different numbers of parameters. Within a family of models, adding parameters generally improves the fit. BMDS reports the Akaike Information Criterion (AIC) to aid in comparing the fit of different models. When comparing the fit of two or more

models to a single data set, the model with the lesser AIC was considered to provide a superior fit. The benchmark response (BMR) level used for this analysis was set at a standard deviation (SD) value of 1.0. This value was chosen based on EPA draft guidelines for BMC analysis (U.S. EPA, 2000), in the absence of a clear biological rationale for selecting an alternative response level.

The following guidance was followed with regard to the choice of the Benchmark Concentration Lower Limit (BMCL) to use as a point of departure for calculation of the pRfC. This guidance is consistent with recommendations in U.S. EPA's BMC guidance (2000). For each endpoint, the following procedure is recommended:

- 1. Models with an unacceptable fit are excluded.
- 2. If the BMCL values for the remaining models for a given endpoint are within a factor of 3, no model dependence is assumed, and the models are considered indistinguishable in the context of the precision of the methods. The models are then ranked according to the AIC, and the model with the lowest AIC is chosen as the basis for the BMCL.
- 3. If the BMCL values are not within a factor of 3, some model dependence is assumed, and the lowest BMCL is selected as a reasonable conservative estimate, unless it is an outlier compared to the results from all of the other models. Note that when outliers are removed, the remaining BMCLs may then be within a factor of 3, and so the criteria given in item 2 would be applied.
- 4. The BMCL values from all modeled endpoints are compared, along with any NOAELs or LOAELs from data sets that were not amenable to modeling, and the lowest NOAEL or BMCL is chosen.

The BMC results are summarized in Table 11 and the individual BMDS model run output is provided in the attachments.

For modeling of the absolute liver weight data set, a constant variance model was appropriate (see test 2 in the BMDS output). The power and polynomial models both defaulted to a linear model. None of these linear models fit the data well. The Hill model provided an excellent fit to the data, as indicated by visual inspection of the fit and the comparison of the maximum likelihood estimates for the fitted model to the optimum model (shown as model A1 in the BMDS output). The linear models failed to provide an adequate fit to the full data set, since they did not accommodate the plateau of the concentration-response curve between the mid- and high-concentrations. BMC modeling was redone using a truncated data set (high concentration group removed) to optimize the fits of these models. Removing the high concentration resulted in good fits for the linear models (the power and polynomial models again defaulted to linear) as indicated by the AIC and goodness-of-fit p-values. The Hill model could not be run with the truncated data set since at least four concentration groups are required to provide a model fit.

Adequate fits to the data were achieved when the high concentration data were removed. An argument could be made for using these results as the best estimate for the data set, since an adequate fit was achieved with fewer parameters than for the Hill model using the full data set. However, the BMCL estimate for the full data set was on the border of 3-fold lower than for the truncated data set, which would suggest that the lower BMCL should be selected. Furthermore, comparison of the chi square residuals in the range of the NOAEL concentration suggests that the Hill model provided a better fit of the data in the low concentration region than the linear models using the truncated data. Finally, since

there was no biological rationale for removing the high concentration data from the modeling, an adequate model fit for the full data set is preferred over the model fit for the truncated data set. Based on these considerations, the BMC of 0.78 mg/m³ and corresponding BMCL of 0.33 mg/m³ are considered the best estimates for the absolute liver weight data set.

The relative liver weight data displayed a similar plateau between the mid- and high-concentration groups. The linear, power, and polynomial models all failed to provide an adequate fit. As for the absolute liver weight data, the Hill model provided an excellent fit to the data, but in this case failed to calculate a BMCL. In the absence of an adequate BMCL estimate for any of the models using the full data set, the data were remodeled with the high concentration group data removed. The power and polynomial models were nearly linear, as indicated by the parameter estimates in the BMDS output. The linear, power, and polynomial models all provided a similar, and very good visual fit to the data. The goodness-of-fit statistic for the linear model was 0.9. Although BMDS did not calculate the goodness-of-fit p-values for the power and polynomial models, inspection of the maximum likelihood estimates for these fitted models as compared to the optimum model (model A1 in the BMDS output) confirmed the good fit. The linear model provided a similar BMC and BMCL estimate as the power and polynomial models, but required less parameters to do so (i.e., as reflected in the lower AIC). Therefore, the BMC of 1.3 mg/m³ and the corresponding BMCL of 0.94 mg/m³ are considered the best estimates for the data set for relative liver weight.

At the time of the meeting the CATT panel did not provide a recommendation on whether absolute or relative liver weight should be considered more appropriate as the critical effect. Both of these measures were significantly increased beginning in the 7.6 mg/m³ study concentration group. One would not expect a difference in the sensitivity of these two measures in this case, because there was no change in body weight (the basis for calculating relative liver weight) at the NOAEL. Therefore, both absolute and relative liver weight changes are considered to be an adequate basis for the critical effect. Based on this consideration, the lower of the BMCL estimates for the absolute and relative liver weight changes is the most appropriate basis for deriving the pRfC. The BMC of 0.78 mg/m³ with the corresponding BMCL of 0.33 mg/m³ for increased absolute liver weight are the best estimates from the BMC modeling results. The BMCL of 0.33 mg/m³ is the most appropriate choice as the critical effect level for derivation of the pRfC, because the BMCL is lower than either the NOAEL of 0.61 mg/m³ for liver effects or the NOAEL of 0.81 mg/m³ for pulmonary effects in this study.

Selection of uncertainty factors

As described in the technical meeting notes, the CATT panel unanimously agreed on the choice of 3 for extrapolation from an animal study (UF_A), a factor of 10 to account for variability in human sensitivity (UF_H), and a factor of 1 for extrapolation from study NOAEL or BMDL (UF_L). The CATT panel considered the selection of U.S. EPA's other two factors, for extrapolation from a study of less-than-lifetime duration (UF_S) and for database insufficiencies (UF_D), to be dependent on whether liver or lung was ultimately selected as the critical effect. The panel was not unanimous in selection of the UFs or UF_D for either organ, but a clear majority vote was obtained for these UFs regarding liver toxicity.

Based on the liver as a critical effect, panel members recommended values of either 1 (one vote), 3 (six votes) or 10 (1 vote) for UF_S, and values of 3 (six votes) or 10 (two votes) for UF_D. Therefore, based on the liver as the critical effect, the composite UF would range from 100 to 1000, depending on the selection of the values for UF_S and UF_D. The majority vote of the CATT panel (Table 5) supported a factor of 3 for UF_S and 3 for UF_D. Based on these values, a composite UF of 300 for liver effects was calculated.

Based on the lung as the critical effect, panel members recommended values of either 1(three votes), 3 (three votes) or 10 (two votes) for UF_S , and values of 1 (one vote), 3 (five votes), and 10 (two votes) for UF_D . Therefore, with the lung as the critical effect the composite UF would range from 30 to 3000. The majority of the CATT panel supported a value of 3 for UF_D based on lung effects. A clear majority vote was not determined for any one value for the UF_S ; however, six votes were cast for a value lower than 10 and five votes were cast for a value higher than one; thus the median value of 3 would be a reasonable choice. Therefore, values of 3 for both UF_D and UF_S for lung effects would also result in a composite UF of 300.

However, it is important to note that the panel could not arrive at a consensus on the overall magnitudes of UF_S and UF_D, because of the numerous uncertainties with the inhalation database. The resulting range in the uncertainty factor was generally considered reasonable by the panel, with values falling within this range being indistinguishable from each other.

Calculation of the pRfC

Liver toxicity was identified as the critical effect because it was more sensitive to C8 than the lung (i.e., liver toxicity had a lower NOAEL or BMCL than lung), the composite UF ranged from 100 to 1000 and was 300 based on the majority vote.

The pRfC is calculated as follows:

```
pRfC (mg/m³) = critical effect level / composite UF pRfC \text{ range} = 0.33 / 1000 = 0.00033 \text{ mg/m³ (or rounded to } 0.3 \text{ µg/m³})to = 0.33 / 100 = 0.0033 \text{ mg/m³ (or rounded } 3.3 \text{ µg/m³})pRfC \text{ (majority vote)} = 0.33 / 300 = 0.0011 \text{ mg/m³ (or rounded to } 1 \text{ µg/m³})
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Therefore, the recommended pRfC based on the majority vote for a composite UF of 300 is 1 microgram per cubic meter of air ($\mu g/m^3$) with a range from 0.3 $\mu g/m^3$ to 3.3 $\mu g/m^3$.

7	able 11. Benchmark Dose M	Iodeling Results for	or C8 ^a				
Model/Data Set	AIC	P-value	BMC ^b	BMCL			
Absolute Liver Weight -All	Data Modeled			·			
Linear	62.58 ^c	<0.001 ^d	31	19			
Hill	48.67	1.0 ^e	0.78	0.33			
Power	62.58°	< 0.001	31	19			
Polynomial	62.58°	< 0.001	31	19			
Absolute Liver Weight - Hig	h Concentration not Modelec	ì					
Linear	38.22°	0.72	1.6	1.1			
Power	38.22°	0.29 ^d	1.6	1.1			
Polynomial	38.22°	0.72	1.6	1.1			
Hill	Insufficient l	Insufficient Number of data points to run model					
Relative Liver Weight – All l	Data Modeled						
Linear	-167.65°	< 0.001	21	15			
Hill	-184.29	1.0 ^e	1.1	Failed			
Power	-167.65°	< 0.001	21	15			
Polynomial	-167.65°	< 0.001	21	15			
Relative Liver Weight - High	Concentration not Modeled	•	•	•			
Linear	-137.04 ^c	0.90	1.3	0.94			
Power	-135.05°	Failed	1.5	0.94			
Polynomial	-135.05°	1.0°	1.5	0.94			
Hill	Insufficient l	Number of data po	ints to run mode	1			

^a Modeling was performed based on absolute and relative liver weight results reported in Kennedy et al. (1986).

^b BMC and BMCL are based on benchmark response of 1SD. Results are presented in units of mg/m³. BMC and BMCL estimates in bold type are the estimates judged to be the best estimates for each endpoint. "Failed" indicates that BMDS was unable to produce the estimate or the information required to be able to present a value.

^c Corrected from erroneous BMDS output. Errors were identified in the degrees of freedom (DF) provided in the output for the fitted model in several cases. For these cases, the AIC was calculated independently using the log likelihoods provided in the output and the correct number of DF. Similarly, the goodness-of-fit p-values were corrected by calculating manually the chi square p-value using the appropriate number of DF.

^d This model provided an identical fit to the linear and polynomial models. The reported P-value reflects a difference in the maximum likelihood estimate for the comparison model (Model A1 in the BMDS output) across the three models. This difference the maximum likelihood estimate should be the same for all three models, since this estimate is model independent.

^e A fit that maximizes the likelihood is assigned a p-value of 1.0, even if there were no degrees of freedom for a formal statistical test. The maximized likelihood is given by model A1 for constant variance models and model A2 for non-constant variance models. Models A1 and A2 are independent of the model chosen to fit the data (e.g., power, polynomial, Hill model) and provide the best match possible to the mean and standard deviation for each dose level.

Calculation of an Air Screening Level

As described in the technical meeting notes, U.S. EPA Region 9 methodology was judged by the CATT panel to be an appropriate basis for deriving the air screening level. The following standard formula was used to calculate the air screening level:

Air Screening Level (
$$\mu$$
g/m³) = THQ x RfDi x BW x AT x 1000
EF x ED x air IR

Note: RfDi (mg/kg-day) = RfC x
$$\frac{20\text{m}^3/\text{d} (IR)}{70 \text{ kg (BW)}}$$

Where:

THQ = Target Hazard Quotient, assumed to be 1

RfDi = The RfC expressed in terms of dose, mg/kg-day RfC = The inhalation reference concentration (mg/m³) BW = Body weight, assumed to be 70 kg for adults

AT = Averaging time, 10,950 days, the exposure duration expressed in days EF = Exposure Frequency, 350 days/year, the average number of days each

year people are exposed

ED = Exposure duration, 30 years, the average number of years people are exposed

IR = Inhalation rate for air screening levels, $20 \text{ m}^3/\text{day}$

Using this equation, the air screening level ranges from $0.3 \mu g/m^3$ to $10 \mu g/m^3$. Using a reasonable median value, the air screening level would be $1.1 \mu g/m^3$ (or rounded to $1 \mu g/m^3$).

2.4 SUMMARY OF FINDINGS

The key studies, critical effects and levels, uncertainty factors, and provisional risk factors developed by the CATT toxicologists are summarized in Table 12.

Tab	le 12. Summary of RfD and Rf	C Values for C8	Determ	nined by	/ the C	ATT 1	Гохісою	ogists	
Reference	Critical Effect	Critical Effect Level ^a	UF₄	UF _∺	UF	UFs	UF₀	Composite UF°	RfD/RfC
Oral Studies									
Palazzolo et al. (1993) ^c 90-day rat study	Increased relative liver weight with histopathology in male rats	0.47 (NOAEL in males) 0.72 (BMDL)	10	10	1	1	1	100	0.005 0.007
York et al. (2002) Two-Generation rat study	Increased liver weight in male rats, supported by histopathology at higher doses (histopathology was not examined at the lowest dose, but incidence of hypertrophy was 100% at next highest dose).	0.42 (BMDL in males) ^a	10	10	1	1	1	100	0.004
RikerLaboratories (1983) Two-year rat study	Hepatic megalocytosis in male rats.	0.73 (BMDL in males)	10	10	1	1	1	100	0.007
Thomford et al. (2001)°26-week cynomolgus monkey study	Decreased thyroid hormone levels in male cynomolgus monkeys, and supported by a NOAEL at the same dose for clinical signs of toxicity in the co-critical rhesus monkey study (Goldenthal et al., 1978b)	3 - 10 (LOAEL in males)	10	10	3	3	1	1000	0.003 - 0.01

Kennedy et al. (1986)	Increased liver weight supported by histopathology	0.61(NOAEL - HEC⊪males)	3	10	1	3	3	300	1
Two-week rat study	and clinical chemistry in male	0.33 (BMCL,							
	rats	BMC 0.78							
		absolute liver weight)							
		0.94(BMCL,							
		BMC 1.3							
		relative liver							
		weight)							
Dermal Studies									
Kennedy et al. (1985)	Increased liver weight in male	4.2° (LOAEL							Data
, ,	rats	in males)							Inadequate
Two-week rat study									

- a. Oral and Dermal effect levels and RfDs are presented in units of mg/kg-day, while the inhalation critical effect level and RfDs are presented in units of mg/m³
- b. Areas of uncertainty addressed by uncertainty factors are: animal to human extrapolation (A); intrahuman variability and protection of sensitive subpopulations (H); extrapolation from a LOAEL to a NOAEL(L); extrapolation from a subchronic to chronic exposure (S); and lack of a complete database (D)
- c. The subchronic study by Goldenthal et al. (1978a) could serve as a supporting study for liver effects in rats.
- d. BMDL is the 95% lower confidence limit on the dose corresponding to a response level of 10% or an increase of 1SD in the continuous endpoint being assessed. Only modeling results that provided the lowest value and provided an adequate fit to the data are provided.
- e. The subchronic study in rhesus monkeys by Goldenthal et al. (1978b) is a co-critical study for clinical signs of toxicity in monkeys.
- f. These studies are not adequate for derivation of an IRIS quality RfD/RfC of even low confidence. The values shown could be used to derive a provisional value. Derivation of the RfC or RfD via route-to-route extrapolation is not supported by the available toxicokinetic data. Consensus on the values for UF_s and UF_o was not reached by the panel; however, a majority vote was obtained for a value of 3 for both these UFs in reference to liver as the target organ. See text of this report for ranges of UFs and SLs based on the range distribution of the votes for UFs.
- g. 4.2 mg/kg-day reflects the study dose of 20 mg/kg adjusted for discontinuous exposure.

I agree that the notes as presented accurately reflect the panel's discussion and conclusions during the May 6-7, 2002 C8 Assessment of Toxicity Toxicologists Panel Meeting, and that the post meeting actions taken to develop the pRfC and Air Screening Level are in accordance with the instructions provided to *TERA* by the panel. (Original signatures are on file at DEP.)

John Cicmanec, D.V.M., M.S., ACLAM, USEPA	ORD Date	
Joan Dollarhide, M.S., M.T.S.C., J.D., TERA	Date	
Michael Dourson, Ph.D., D.A.B.T., TERA	Date	
Gerald Kennedy, DuPont	Date	
Andrew Maier, Ph.D., C.I.H., TERA	Date	
Samuel Rotenberg, Ph.D., USEPA Region 3	Date	
Jennifer Seed, Ph.D., USEPA Headquarters OPPT	Date	
Dee Ann Staats, Ph.D., DEP (Chairperson)	Date	
John Wheeler, Ph.D., D.A.B.T., ATSDR	Date	
John Whysner, M.D., Ph.D., D.A.B.T.	Date	

3. 0 COMPARISON OF SCREENING LEVELS TO SITE-RELATED DATA

After the SLs for air, water, and soil were determined, DEP compared these SLs to the site-related data that has been collected to date. These comparisons are summarized below. The work of the CATT was only one facet of an investigation that continues beyond the issuance of this report. The GIST is expected to issue a report of the groundwater and surface water data in early 2003. The air modeling effort continues and is currently focusing on determining the results of the air emissions reduction efforts by DuPont required in the consent order as a 50% reduction in overall emissions (both air and water) by the end of 2003. Upgrades were completed in June 2002 which included the installation of a new scrubber and increased height of the primary C8 emissions stack.

Water

To date, of the 188 samples collected from private wells, cisterns, and springs, 50 were used for drinking water and none exceeded the 150 ppb health protective water SL for C8. Also to date, nine public water supply facilities in West Virginia have been analyzed for C8, including Belleville Locks and Dam, Blennerhassett Island, General Electric, Lubeck Public Service District (PSD), Mason County PSD, Parkersburg PSD, Racine Locks and Dam, New Haven Water Department, and Ravenswood. None of the drinking water from these facilities contained concentrations of C8 that exceeded the 150 ppb water SL. In fact, the concentrations of C8 in public water supplies were all below 2 ppb, below 15 ppb in private non-drinking water, and below 3 ppb in private drinking water wells in West Virginia. Samples were collected from Ohio public and private water supplies. Although C8 levels in some Ohio private water supplies were higher than those detected in West Virginia, none of these samples contained C8 concentrations above the water SL. These data have been provided to Ohio EPA and DEP will continue to share information with throughout the remainder of this investigation. The DEP notes that the water SL is higher than DuPont's internal community exposure guidelines for drinking water of 1 or 3 ppb; however, these guidelines were developed in the early 1990s and based solely on a two-week inhalation study from 1986. Since then significant additional toxicological data have been collected and the CATT water SL is based on a comprehensive examination of all available information. Sampling of the Ohio River has begun; preliminary analytical results are expected from the laboratory in September 2002. To date, no analysis has been performed to measure C8 in soils in West Virginia on private property; therefore, no comparison can be made to the soil SL.

Air

Mathematical computer models that incorporate weather conditions, chemical characteristics, and facility measurements were utilized by DEP to simulate the ambient air concentrations of C8. Based on actual emissions data from the DuPont WW facility for the year 2000, the DEP modeling efforts predicted a maximum C8 concentration in air of approximately $2.7 \,\mu\text{g/m}^3$ at the facility fence line along the Ohio River. The maximum modeled C8 air concentration in the West Virginia residential area adjacent to the facility was approximately $0.2 \,\mu\text{g/m}^3$ annual average. Predicted C8 air concentrations across the Ohio River from the WW facility in Ohio residential areas were greater than those predicted in residential areas in West Virginia. These data have been provided to Ohio EPA and DEP will continue to share information with Ohio EPA throughout the remainder of this investigation. Results of similar subsequent air modeling efforts conducted by DuPont are consistent with those of the DEP. Air modeling information can be obtained from the DEP Division of Air Quality.

The DEP's Divisions of Water Resources and Air Quality are currently reviewing all relevant air and water data to determine DuPont's compliance with the November 2001 consent order between DEP and DuPont.

To: Bertrand, Charlotte[Bertrand.Charlotte@epa.gov]

From: Dourson, Michael

Sent: Fri 11/3/2017 2:52:32 PM

Subject: Re: Travel to Conference in Saudi Arabia

Ok

Sent from my iPhone

On Nov 3, 2017, at 10:51 AM, Bertrand, Charlotte < Bertrand. Charlotte@epa.gov > wrote:

We will certainly ask for a follow-up seminar. You'll see in my next message that State would like to see us attend, so I have given my ok. I've asked Kaitlin to set up a meeting with the scientist, Rick and the IO today to discuss the presentation, talking points and what we'd like to get out of the attendance. Will include you.

From: Dourson, Michael

Sent: Friday, November 03, 2017 10:43 AM

To: Bertrand, Charlotte < <u>Bertrand.Charlotte@epa.gov</u>> **Subject:** Re: Travel to Conference in Saudi Arabia

Charlotte

This for the opportunity to help. In general, I favor these kind of interactions. However, the overall meeting is quite broad. What does our scientist expect to get out of this meeting. Is s/he willing to give us all a seminar on findings afterwards?

Michael

Sent from my iPhone

On Nov 3, 2017, at 9:30 AM, Bertrand, Charlotte < Bertrand. Charlotte@epa.gov > wrote:

Thanks for offering your thoughts on the science benefits of attendance. Invite and link to conference is below.

Sent from my iPhone

Begin forwarded message:

From: "Bertrand, Charlotte" < Bertrand. Charlotte@epa.gov >

Date: November 2, 2017 at 6:25:30 PM EDT

To: "Dieu, Martin" < Dieu. Martin@epa.gov>, "Kasman, Mark"

< <u>Kasman.Mark@epa.gov</u>>

Cc: "Nishida, Jane" < Nishida.Jane@epa.gov>, "Beck, Nancy"

< beck.nancy@epa.gov>, "Wise, Louise (Wise.Louise@epa.gov)"

< Wise. Louise@epa.gov>

Subject: Travel to Conference in Saudi Arabia

Martin and Mark -

Ex. 5 - Deliberative Process

https://sfdaconf.com/en/index.php

Thank you for your assistance,

Charlotte

Charlotte Bertrand

Acting Principal Deputy Assistant Administrator

Office of Chemical Safety and Pollution Prevention Phone (202) 564-2910

<Invitational letter from SFDA - Alaa Kamel.pdf>

To: Bowman, Liz[Bowman.Liz@epa.gov]

Cc: Jackson, Ryan[jackson.ryan@epa.gov]; Beck, Nancy[Beck.Nancy@epa.gov]

From: Dourson, Michael

Sent: Tue 10/31/2017 4:13:43 PM

Subject: Re: RE:

Ryan

No worries. I am in a meeting now but can come over. Just say the word...

Michael

Sent from my iPad

On Oct 31, 2017, at 12:11 PM, Bowman, Liz < Bowman.Liz@epa.gov > wrote:

Yes, send it over...I have an hour until the pen and pad

From: Jackson, Ryan

Sent: Tuesday, October 31, 2017 12:07 PM

To: Beck, Nancy < Beck. Nancy@epa.gov >; Dourson, Michael

dourson.michael@epa.gov>; Bowman, Liz Bowman, Liz Bowman, Liz <a href

Subject:

I need Ex. 5 - Deliberative Process

Ryan Jackson

Chief of Staff

U.S. Environmental Protection Agency

Ex. 6 - Personal Privacy

To: Cc: From: Sent: Subject:	Griffo, Shannon[Griffo.Shannon@epa.gov] Fugh, Justina[Fugh.Justina@epa.gov] Dourson, Michael Tue 11/28/2017 11:49:23 PM RE: Request from OGC/Ethics
Shannon	
be contac	act information for Dr. Andrew Maier is: maierma@ucmail.uc.edu . I believe Ellen can ted at ellen.rozenson@uc.edu , but am not sure of this. My previous email contact list is UC computer, which, of course, I do not now have.
Cheers!	
Michael	
Sent: Tue To: Dour Cc: Fugh	riffo, Shannon esday, November 28, 2017 2:26 PM rson, Michael <dourson.michael@epa.gov> , Justina <fugh.justina@epa.gov> RE: Request from OGC/Ethics</fugh.justina@epa.gov></dourson.michael@epa.gov>
Hi Micha	el,
Please se	nd me the contact information for Dr. Maier and Ms. Rozenson.
Thanks in	n advance,
Shannon	

Ethics Attorney
Office of General Counsel, Ethics
U.S. Environmental Protection Agency
(202) 564-7061
Griffo.Shannon@epa.gov
From: Dourson, Michael Sent: Monday, November 20, 2017 1:00 PM To: Griffo, Shannon < Griffo. Shannon@epa.gov> Cc: Fugh, Justina < Fugh. Justina@epa.gov> Subject: RE: Request from OGC/Ethics
Shannon
No worries, but I may not be able to respond to your request. Although I was the lead of the TERA Center for the first year, and the senior scientist for the Risk Science Center for 1 year (after two centers combined), I am not allowed to give this information out routinely. I can answer general questions, of course, but for specific information you may need to contact either Dr. Andrew Maier, the RSC Director, or Ms. Ellen Rozenson, the Department Financial lead.
Please let me know if you need their contact information.
Cheers!
Michael

Shannon Griffo

... L. Dourson, PhD., DABT, FATS, FSRA Senior Advisor to the Administrator U.S. Environmental Protection Agency dourson.michael@epa.gov 202-564-2463 www.epa.gov From: Griffo, Shannon Sent: Monday, November 20, 2017 12:25 PM To: Dourson, Michael < dourson.michael@epa.gov> Cc: Fugh, Justina < Fugh. Justina@epa.gov> Subject: Request from OGC/Ethics Michael, We're sorry that we couldn't connect last week. Justina wasn't in the office on Friday (or today). And I think you are gone the rest of this week. We'll try again once we are all back after the holiday. In the meantime, we wanted to give you a heads up that we need to know what University of Cincinnati contracts you worked on over the past two years, for whom and for what, and we'd like to see the scopes of work for these contracts if possible. When we meet, we'll provide more background and information into this request. Have a Happy Thanksgiving! Thanks,

Shannon

Shannon Griffo

Ethics Attorney

Office of General Counsel, Ethics

U.S. Environmental Protection Agency

(202) 564-7061

Griffo.Shannon@epa.gov

To: Dinkins, Darlene[Dinkins.Darlene@epa.gov]

From: Dourson, Michael

Sent: Tue 11/21/2017 8:52:26 PM Subject: RE: OPP General Agenda

Thanks!

From: Dinkins, Darlene

Sent: Tuesday, November 21, 2017 3:07 PM

To: Dourson, Michael <dourson.michael@epa.gov>

Subject: FW: OPP General Agenda

Hi Mike,

Please find attached the agenda for the OPP General this afternoon.

Call-in # Ex. 6 - Personal Privacy

Darlene Dinkins

Office of Pesticide Programs

U.S. Environmental Protection Agency

(703) 305-5214

To: Fugh, Justina[Fugh.Justina@epa.gov]; Hanley, Mary[Hanley.Mary@epa.gov]

Cc: Griffo, Shannon[Griffo.Shannon@epa.gov]

From: Dourson, Michael

Sent: Wed 11/15/2017 1:03:39 AM

Subject: RE: For Review: Draft Response to SEPW Minority Letter

mdFINAL Dourson QFRs 11.14.2017..docx

mdDourson Response to SEPW Minority Letter.draft 11.14.17..docx

Justina and Mary

Please find attached my strike and replace for the answers, and also a tweaked letter (which I did not do as S&R---sorry). I would be more than happy to take additional guidance. My take on the questions on page 3 of the incoming letter were that they reflected summaries of the prior questions. Thus, I gave what are probably verbose answers that should reflect an overview of the prior answers. I will defer to your collective wisdom on how best to respond to these questions, however, or to leave them as unresponded.

Cheers!

Michael

From: Fugh, Justina

Sent: Tuesday, November 14, 2017 6:20 PM **To:** Hanley, Mary <Hanley.Mary@epa.gov>

Cc: Dourson, Michael <dourson.michael@epa.gov>; Griffo, Shannon

<Griffo.Shannon@epa.gov>

Subject: RE: For Review: Draft Response to SEPW Minority Letter

Yes, we will look at this ASAP. I'll make edits tonight and then ask Shannon Griffo to look it over tomorrow. She drafted a recusal statement but I have not yet sent it forward. I'll email Shannon tonight with an update so she'll have the latest. I'm going to be out of pocket most of the day tomorrow, so you can chat with Shannon at 564 7061.

Justina Fugh | Senior Counsel for Ethics | Office of General Counsel | US EPA | Mail Code 2311A | Room 4308 North, William Jefferson Clinton Federal Building | Washington, DC 20460 (for ground deliveries, use 20004 for the zip code) | phone 202-564-1786 | fax 202-564-1772

From: Hanley, Mary

Sent: Tuesday, November 14, 2017 5:00 PM **To:** Fugh, Justina < Fugh.Justina@epa.gov >

Cc: Dourson, Michael < dourson.michael@epa.gov>

Subject: FW: For Review: Draft Response to SEPW Minority Letter

Hello Justina,

We are finalizing a response to the SEPW Minority letter in which they request that Mike answer some questions (page 1-2 on the incoming letter attached) and also update his earlier answers to some of the original QFRs. I have flagged a few items for you and Mike in the Dourson response letter. Would you be able to help with those items? We will need to finalize the pieces and get them to OCIR on Thursday. Many thanks. For more information on the attachments, etc. see my email below. Also, once I have edits from OCSPP I will share the next version with Brian Grant and company sometime tomorrow afternoon. Thanks.

Cheers

Mary

From: Hanley, Mary

Sent: Tuesday, November 14, 2017 4:47 PM

To: Beck, Nancy <beck.nancy@epa.gov>; Bertrand, Charlotte <Bertrand.Charlotte@epa.gov>;

Louise Wise (Wise.Louise@epa.gov) < Wise.Louise@epa.gov>; Michael Dourson

(dourson.michael@epa.gov) <dourson.michael@epa.gov>

Cc: Kaitlin Keller (keller.kaitlin@epa.gov) <keller.kaitlin@epa.gov>
Subject: For Review: Draft Response to SEPW Minority Letter

Hello,

Attached is the draft letter response and an attachment with the suggested updated answers to a group of the original QFRs that were identified in the incoming letter. Changes to the answers are shown in redline. Also attached is the incoming letter and another copy with Mike's initial draft answers embedded. I am also including the Bodine response letter which we are using a model for responding.

Please note that draft letter response only addresses the questions presented on pages 1-2 of the letter. Rather than attempt to answer the comments on page 3 of the incoming letter we are responding to the specific QFRs in the attachment. The comments on page 3 of the incoming reference QFRs but it is impossible to pin down which QFRs they refer to. As you review the QFRs please keep in mind the comments on page 3 to see if any updates might address the comments as appropriate.

Note the QFR attachment needs some formatting so don't worry about the page numbering, etc. That will be taken care of in the next version. I will wait for your edits before sending to OGC for their review. The only exception is that I will send the response letter to Justine to see if she can assist with the items I flagged for her/Mike.

Please let me know of any questions.

Cheers

Mary

To: Beck, Nancy[Beck.Nancy@epa.gov]

From: Dourson, Michael

Sent: Mon 10/23/2017 12:56:19 PM

Subject: Re:

Hmm... thanks

Sent from my iPhone

On Oct 23, 2017, at 8:26 AM, Beck, Nancy < Beck. Nancy@epa.gov > wrote:

No worries Mike. Those programs cannot be added to EPA iPads, essentially making the iPad useful only for email. You will not be able to access any of your files on the iPads either.

Ex. 6 - Personal Privacy

Yes, the 1pm can be covered by Louise today. The Administrator does not attend that meeting.

Nancy B. Beck, Ph.D., DABT Deputy Assistant Administrator, OCSPP

P: 202-564-1273

M: Ex. 6 - Personal Privacy

Beck.Nancy@epa.gov

On Oct 23, 2017, at 6:05 AM, Dourson, Michael dourson.michael@epa.gov> wrote:

Nancy

Sorry for the confusion. I will ask IT to add word, excel, and PP to my ipad. This will allow more clear commentaries.

I am over at PY during Pruitt's 1 pm senior staff meeting. You have this covered, correct?

Cheers!

Mike

Sent from my iPad

On Oct 22, 2017, at 10:51 PM, Beck, Nancy < Beck. Nancy@epa.gov > wrote:

Not sure what you mean by interface but generally RAD is supportive of using high throughput method.

Nancy B. Beck, Ph.D., DABT
Deputy Assistant Administrator, OCSPP
P: 202-564-1273
M: Ex. 6 - Personal Privacy
Beck. Nancy@epa.gov

On Oct 21, 2017, at 10:51 AM, Dourson, Michael dourson.michael@epa.gov wrote:

Interface: Nancy, has ORD been working with our OCSPP colleagues on this interface? If so, what are the thoughts of our RAD folks?

Sent from my iPad

Subject: Epidemiology studies Dear Nancy and Charlotte It would be very helpful to receive a copy of all relevant epi studies on chlorpyrifos prior to our Ex. 5 - Deliberative Process briefing next week. Also, I am thinking Ex. 5 - Deliberative Process From an administrator's point of view, Ex. 5 - Deliberative Process Ex. 5 - Deliberative Process Thanks! Michael... ... L. Dourson, PhD., DABT, FATS, FSRA Senior Advisor to the Administrator U.S. Environmental Protection Agency dourson.michael@epa.gov 202-564-2463 www.epa.gov

Beck, Nancy[beck.nancy@epa.gov]; Bertrand, Charlotte[Bertrand.Charlotte@epa.gov]

To:

From:

Sent:

Dourson, Michael

Fri 11/10/2017 4:10:37 PM

To: Beck, Nancy[beck.nancy@epa.gov]

From: Dourson, Michael

Sent: Tue 11/21/2017 7:27:17 PM

Subject: RE: OPP Acute Dietary Exposure: 99.9th Percentile

Thanks!

----Original Appointment-----

From: Keller, Kaitlin On Behalf Of Beck, Nancy Sent: Tuesday, November 21, 2017 1:56 PM

To: Bertrand, Charlotte; Dourson, Michael; Dawson, Jeffrey; Vogel, Dana; Lowe, Kelly;

Van Alstine, Julie; Perlis, Robert; Lowit, Anna; Keigwin, Richard

Cc: Wilbur, Donald; Dyner, Mark; Huskey, Angela; Wise, Louise; Hughes, Hayley; Davis,

Donna

Subject: OPP Acute Dietary Exposure: 99.9th Percentile

When: Tuesday, November 21, 2017 2:15 PM-3:00 PM (UTC-05:00) Eastern Time (US &

Canada)

Where: S-12100 PYS; (Call in number Ex. 6 - Personal Privacy

To: Keller, Kaitlin[keller.kaitlin@epa.gov]; Bertrand, Charlotte[Bertrand.Charlotte@epa.gov]; Beck,

Nancy[beck.nancy@epa.gov]

Cc: Bolen, Derrick[bolen.derrick@epa.gov]

From: Dourson, Michael

Sent: Mon 11/13/2017 11:49:12 PM
Subject: RE: BEAD lab Friday 11/17 logistics

Kaitlin

This is not a problem with me. I will be flying out of RR airport later that evening anyway.

Cheers!

Michael

From: Keller, Kaitlin

Sent: Monday, November 13, 2017 6:25 PM

To: Bertrand, Charlotte <Bertrand.Charlotte@epa.gov>; Beck, Nancy <Beck.Nancy@epa.gov>;

Dourson, Michael dourson, Michael dourson.michael@epa.gov)

Hello—BEAD is looking forward to your trip to the Fort Meade lab this Friday and is working out the agenda, but I first have a couple of questions on logistics for you, with a detailed breakdown below. In a nutshell, the question is can you plan to start and end your day at Potomac Yards on Friday?

- 1) Would you be able to meet at 7:45am at Potomac Yards? Unfortunately, the motor pool can't be scheduled that early, so you'd have to plan to start your day there via metro or parking (see #2). However, if you cannot, please let me know and we will work out another option.
- a. Some explanatory details—The lab is about 30 miles away, and for cars to get onto the Fort Meade campus, drivers have to have a certain badge, (so motor pool is not an option).

Wynne Miller and Anita Pease (BEAD Director and Deputy Director) have badges and will drive you in two cars. The tour will start at 9am, and traffic in and out of DC is a consideration to be mindful of.

- 2) Would it be okay for you to go straight back to PY after the tour then call in to the 1pm problem formulation meeting from a conference room there? Since most of the afternoon is at PY, I thought just heading back there would make sense, and that might make starting there easier as well if you'd like to drive.
- a. The tour ends by 12pm, though again, traffic is always a factor on Fridays on the parkway, and you have a 1pm biweekly problem formulation meeting in WJC-E, followed by two meetings in PY--2-4:00pm on chlorpyrifos and 4-4:30pm on WPS. Charlotte-you probably know parking at PY better than me—but I believe you can still do daily parking in PY. There's also a slightly cheaper option just around the corner.
- b. Note: There are some lunch options near PY, and I can book separate conference rooms for work space if anyone is not attending the 1pm problem formulation meeting.

Please let me know if this set up	would wo	rk for you.	We can	also	discuss	this in	the	Weds.
Morning huddle if that's easier.								

Thanks,

Kaitlin

Kaitlin Keller, Special Assistant

Office of Chemical Safety and Pollution Prevention

U.S. Environmental Protection Agency

(202) 564-7098

To: Keigwin, Richard[Keigwin.Richard@epa.gov]; Beck, Nancy[beck.nancy@epa.gov]; Bertrand,

Charlotte[Bertrand.Charlotte@epa.gov]; Wise, Louise[Wise.Louise@epa.gov]

Cc: Sands, Jeffrey[sands.jeffrey@epa.gov]

From: Dourson, Michael

Sent: Wed 11/15/2017 12:24:09 AM

Subject: RE: Request for additional information

Rick

Oh well, they did not seem to like either of our suggestions.

Cheers!

Mike

From: Keigwin, Richard

Sent: Tuesday, November 14, 2017 6:55 PM

To: Dourson, Michael <dourson.michael@epa.gov>; Beck, Nancy <Beck.Nancy@epa.gov>; Bertrand, Charlotte <Bertrand.Charlotte@epa.gov>; Wise, Louise <Wise.Louise@epa.gov>

Subject: Fwd: Request for additional information

FYI

Rick Keigwin

Director, Office of Pesticide Programs

U.S. Environmental Protection Agency

Phone: 703-305-7090

Website: www.epa.gov/pesticides

Sent from my iPhone

Begin forwarded message:

From: "Echeverria, Marietta" < Echeverria. Marietta @epa.gov >

Date: November 14, 2017 at 6:33:47 PM EST

To: "Anderson, Brian" < <u>Anderson.Brian@epa.gov</u>>, "Keigwin, Richard" < <u>Keigwin.Richard@epa.gov</u>>, "Dyner, Mark" < <u>dyner.mark@epa.gov</u>>, "Guilaran, Yu-Ting" < <u>Guilaran.Yu-Ting@epa.gov</u>>

Subject: Fwd: Request for additional information

Sent from my iPhone

Begin forwarded message:

From: "Frazer, Gary" < gary_frazer@fws.gov> Date: November 14, 2017 at 6:25:44 PM EST

To: "Echeverria, Marietta" < echeverria.marietta@epa.gov>

Cc: Gina Shultz < Gina Shultz@fws.gov >, Craig Aubrey < craig aubrey@fws.gov >,

Patrice Ashfield < <u>patrice_ashfield@fws.gov</u> > **Subject: Request for additional information**

Attached please find our request for additional information necessary to complete formal consultation on the effects of reregistering chlorpyrifos, malathion, and diazinon. A hard copy will follow.

We look forward to continuing our work together on this consultation process. Please contact me or Gina Shultz if you have any questions.

Gary Frazer

Assistant Director -- Ecological Services

U.S. Fish and Wildlife Service

(202) 208-4646

To: From: Sent: Subject:	Hanley, Mary[Hanley.Mary@epa.gov] Dourson, Michael Thur 12/7/2017 8:39:08 PM Power POint presentation
Mary	
	A have a template for its power point presentations? Like Nancy, I will be giving a talk eek's SRA meeting and would like to use our standard format.
Thanks!	
Michael.	
L. Do	urson, PhD., DABT, FATS, FSRA
Senior A	dvisor to the Administrator
U.S. Env	ironmental Protection Agency
dourson.1	michael@epa.gov
202-564-	2463
www.epa	<u>.gov</u>

To: Keller, Kaitlin[keller.kaitlin@epa.gov]

From: Dourson, Michael

Sent: Mon 11/13/2017 11:43:40 PM
Subject: RE: OPP EFED All Hands tomorrow

Kaitlin

Well, if it is only a few lines then what about:

• College of Medicine in 1980, followed by board certification in 1985. He received fellowships in the Academy of Toxicological Sciences and the Society for Risk Analysis in 2007 and 2009. He has had the good fortune to work at EPA for 15 years, primarily in ORD, for 21 years at the nonprofit organization, Toxicology Excellence for Risk Assessment, and for 2 years back at his alma mater as a Professor at the Risk Science Center. In addition to publishing over 150 papers with colleagues, he has also been invited to give numerous presentations and chair scores of scientific meetings. However, perhaps his greatest contribution to environmental protection has been in the development of risk assessment skills of numerous colleagues who now contribute their time and talent in a variety of organizations.

Please feel free to modify this text for clarity.

Cheers!

Michael

From: Keller, Kaitlin

Sent: Monday, November 13, 2017 5:55 PM

To: Dourson, Michael dourson.michael@epa.gov **Subject:** RE: OPP EFED All Hands tomorrow

Sorry, I didn't mean to create work for you! I think a few lines would be fine, Marietta would just like to properly introduce you.

I'll have print outs for you in the morning, but I've attached EFED's FY17 accomplishments and
FY18 workplan if you'd like to get an idea of the work they do. Also the agenda is below. Note
that they've added you and Charlotte for intro remarks and Q&A, please let me know if you have
any concerns.

Thanks,

Kaitlin

EFED All Hands Agenda:

- • • • Introductory remarks (Marietta)
- • • • Remarks from Charlotte Bertrand, Acting Principle Deputy Assistant Administrator (10:00 AM)
- • • Remarks from Michael Dourson, Senior Advisor to the Administrator (tentative)
- o Q & A with Charlotte and Michael

- o Q & A with Rick
- • • CFC (Sheena Moore 11:30 AM)
- •====== Q & A

From: Dourson, Michael

Sent: Monday, November 13, 2017 5:50 PM
To: Keller, Kaitlin keller.kaitlin@epa.gov
Subject: RE: OPP EFED All Hands tomorrow

Kaitlin

Yes, I am planning to go over with Charlotte tomorrow. I will have to create a new biographical sketch later this evening!
Cheers!
Michael
From: Keller, Kaitlin Sent: Monday, November 13, 2017 5:31 PM To: Dourson, Michael < dourson.michael@epa.gov> Subject: OPP EFED All Hands tomorrow
Hello Mike,
Will you be able to head over to Potomac Yard tomorrow with Charlotte for the Environmental Fate and Effects Division all hands at 10am? The intent was to pop in for the first half hour as a quick meet and greet. If so, could you send me your bio if you have one handy so that Marietta can introduce you? (Either way, it might be good to have a version for my reference.)
Thanks,
Kaitlin
Kaitlin Keller, Special Assistant
Office of Chemical Safety and Pollution Prevention
U.S. Environmental Protection Agency (202) 564-7098
(202) 00

To: Kovner, Karissa[Kovner.Karissa@epa.gov]

Cc: Bertrand, Charlotte[Bertrand.Charlotte@epa.gov]; Beck, Nancy[beck.nancy@epa.gov]; Wise,

Louise[Wise.Louise@epa.gov] From: Dourson, Michael

Sent: Wed 12/20/2017 1:59:44 PM

Subject: RE: Egyptian Training Request to Dourson

Karissa

Ex. 5 - Deliberative Process

Of course, my portfolio is a bit empty of late, so maybe I am speaking from this perspective!

Cheers!

Mike

From: Kovner, Karissa

Sent: Wednesday, December 20, 2017 8:52 AM **To:** Dourson, Michael <dourson.michael@epa.gov>

Cc: Bertrand, Charlotte <Bertrand.Charlotte@epa.gov>; Beck, Nancy <Beck.Nancy@epa.gov>;

Wise, Louise < Wise.Louise@epa.gov>

Subject: Re: Egyptian Training Request to Dourson

Mike,

Thanks and sorry to come back on this, but in re-reading your response, I realized my email might not have been clear enough.

ORD has indeed offered to facilitate two days, the one day of basic training and the one day at Ft. Meade. Are you thinking maybe asking ORD to extend the basic training to two days instead? That would mean the overall training goes to four days instead of three (2 ORD basic training, 1 ORD Ft. Meade, 1 OCSPP and others); I could circle back with Abdel on that if that's the desire.

Thanks,

Karissa

On Dec 18, 2017, at 5:38 PM, Dourson, Michael < dourson.michael@epa.gov > wrote:

Karissa

Ex. 5 - Deliberative Process

accommodate nis request.

Cheers!

Michael

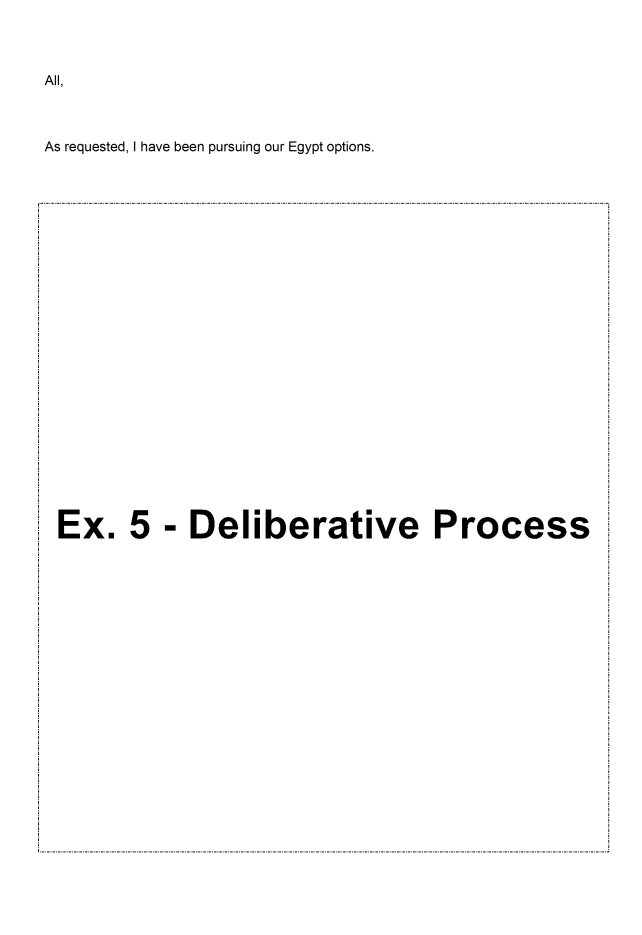
From: Kovner, Karissa

Sent: Monday, December 18, 2017 5:29 PM

To: Bertrand, Charlotte < Bertrand. Charlotte@epa.gov >

Cc: Beck, Nancy < Beck.Nancy@epa.gov >; Wise, Louise < Wise.Louise@epa.gov >;

Dourson, Michael <<u>dourson.michael@epa.gov</u>> **Subject:** RE: Egyptian Training Request to Dourson



Ex. 5 - Deliberative Process

Just let me know what you all, plus Rick and Jeff want, and we can go from there. Thanks, Karissa From: Kovner, Karissa Sent: Monday, November 27, 2017 9:30 PM To: Bertrand, Charlotte < Bertrand. Charlotte@epa.gov > Cc: Beck, Nancy <Beck.Nancy@epa.gov>; Wise, Louise <Wise.Louise@epa.gov> Subject: Re: Egyptian Training Request to Dourson Charlotte, Per our discussion this evening, this is your reminder to send a "thanks Salah, it'll take us a little bit of time to think about this" email in the next day or so to acknowledge his incoming email and potentially begin to lower expectations. I will also reach out to Keigwin as you asked, schedule a meeting for us all with Mike to discuss, and think about some other (smaller, less resource intensive) options for consideration.

Thanks,

Ex. 5 - Deliberative Process

I assume you would like to respond, just let me know if you need anything else from me We would also need to identify a little bit of funding to host a dinner or something.

Thanks, hope this helps. Happy to discuss further if need be.

Karissa

Begin forwarded message:

From: Salah Soliman < Salah Soliman@bibalex.org >

Date: November 21, 2017 at 7:27:00 AM EST

To: "Bertrand.Charlotte@epa.gov" < Bertrand.Charlotte@epa.gov>,

"Kovner.Karissa@epa.gov" < Kovner.Karissa@epa.gov>

Cc: "Bertrand, Charlotte" < Bertrand. Charlotte@epa.gov >, "Beck, Nancy"

<<u>Beck.Nancy@epa.gov</u>>, "Keigwin, Richard" <<u>Keigwin.Richard@epa.gov</u>>,

"Kovner, Karissa" < Kovner. Karissa@epa.gov >, "Dourson, Michael"

<dourson.michael@epa.gov>

Subject: Training Egipians by the USEPA

Dear Charllotte and Karissa,

Hope this message finds you well. Yes it was short but in fact it was very remarkable seeing you Charlotte that evening of Friday, November 8 while in

Crystal City. I am also pleased that you show interest in discussing our proposal requesting the help from your great institution, the USEPA, to train a group of Egyptians on risk evaluation, management and regulations in order to eliminate, reduce and/or minimize stocks and emissions of persistent organic pollutants, POPs.

As I mentioned, while in the States last week, we look forward to your kind support and help in training, for a short time (4-5 days), of about 8 Egyptians representing different Egyptian sectors including the Ministries of Environment, Electricity and Energy, and Agriculture and Land Reclamation. We kindly request the training be conducted in the USEPA facilities by your highly respected experts in order to prepare our trainees to take the lead in directing Egypt, first to assure better and safe environment and second to facilitate its international trade especially with the great USA. This training will very much reflects on safety of all commodities being imported from Egypt by the USA and give more credits to goods being imported by Egypt from the States. Not to mention the impacts for better global environment.

About 7 years ago, with the great help from the USEPA, I initiated a collaborative training program between USEPA and Egypt to train a number of Egyptian analytical chemists on new methods of pesticide residue analysis. The training was conducted in the USEPA Laboratory located in Fort Mead, MD, and lasted for a period of a year. Each group received that training for 6 month on the base that trainees trained for 3 months and then work freely at that Lab for another 3 months. Unfortunately, this training program was terminated because of the uprising of January 25, 2011 occurred in Egypt. I can assure you that the trained Egyptians gained unlimited confidence and on them our two major laboratories of pesticide residue analysis now depend. They clearly reflect to others, not only in Egypt but also in the the whole region, how great to be trained and educated by the USEPA staff members

I am sure that the new proposal will have even much more greater impacts on USA/Egypt relations and deepening the ties between people of the two great nations.

The objectives of this new training proposal can be formulated to cover many POPs related issues. Of these, just for examples, are:

- 1- How EPA sets regulations and guidances to control and or prevent releases of pesticidal POPs, PCBs and the new industrial POPs such as the poly brominated hydrocarbons used as flame retardants and implementation of the Basel and Stockholm Conventions,
- 2- Guidances and regulations to protect, monitor and clean sites potentially contaminated with these pollutants.
- 3- Guidances and regulations to monitor, eliminate and/or minimize releases of unintentional POPs such as dioxins and furans.
- 4- Guidances and regulations to prevent goods with unacceptable levels of POPs residues from entering the USA ports,

- 5- The USEPA recommendations and action plans to safely destructing/eliminating/reducing POPs and POPs highly contaminated goods and articles,
- 6- Site characterization and sampling for pesticides, transformer oil contaminated with PCBs and POPs contaminated areas, and
- 7- Guidances and regulations to assure sustainable management POPs including PCBs and new industrial POPs.

The final contents of the proposed training can be set and agreed upon later. I will be able to come visit you again during December, 2017 or January, 2018 to share with you our needs and help in tailoring the final 4-5 days long training program once you kindly accept this proposal.

The cost of travel, accommodations and local transportations of our trainees, Cairo to Cairo will be covered from our side via the budget of our "Sustainable Management of POPs" Project which is financed by the Egyptian Government and GEF and managed by the WB.

We hope that the training and the training materials and certificate of attendance be as a great complement offered by the USEPA.

I proposed that this training may takes place in USEPA facilities during April or May, 2018.

Once received your kind approval, I will send you names and copies of passports of our nominee to initiate visa requirements.

I am waiting for your kind reply and thank you so much again for making me always proud of have being had the opportunity working with this great agency, the USEPA, since 1974.

With my kindest regards, as ever,

Salah

Salah A. Soliman
Professor of Pesticide Chemistry &Toxicology
Alexandria University
Senior Expert, Bibliotheca Alexandrina
El-Shatby, Alexandria 21526, EGYPT
E-mail: Salah.Soliman@bibalex.org

Ex. 6 - Personal Privacy
Website: www.bibalex.org

www.bibalex.org/yesbu

Sent from my iPad

From: Dourson, Michael [dourson.michael@epa.gov]

Sent: Monday, November 13, 2017 5:49 PM

To: Salah Soliman

Cc: Riham AbdelHamid; Bertrand, Charlotte; Beck, Nancy; Keigwin, Richard;

Kovner, Karissa

Subject: Re: Thank you for your time and generosity

Salah

Well, the delight was with myself and EPA colleagues who had the benefit of your lecture on pesticide use in Egypt, and discussions before and afterwards on ways to improve the assessment of risk and management of pesticides for increased food production. I also agree with you that additional conversations over dinner were insightful, and thoroughly enjoyed them.

Charlotte was most impressed with the short time she spoke with you and Mohammed. As the lead in the Office of Chemical Safety and Pollution Prevention, she will be most interested in potential training opportunities. I also wish to give you our contact for POPs and PICs. She is Karissa Kovner. Please feel free to contact either of these gentle people. Both of them are copied on this email.

Cheers!

Michael...
...L. Dourson
Senior Advisor to the Administrator
US EPA
Washington, DC

Sent from my iPad

On Nov 13, 2017, at 9:43 AM, Salah Soliman <<u>Salah.Soliman@bibalex.org</u>> wrote:

Dear Michael,

This message should have sent to you on Friday evening. Can't tell how

much that I am happy that I had the chance seeing you in Crystal City and touched and delighted to have had such marvelous dinner and outstanding discussion with you that evening.

I am writing this right after arriving into Cairo Airport while riding to Alexandria.

Our meeting at the USEPA building and the time you shared with us while there will indeed positively reflects on cooperation to strengthen the relations between our institutions and countries.

Tomorrow, I will send you and Charolette more details and an official request regarding the training I mention then. I appreciate that you kindly send me her email.

I wish you all the best with my kindest regards, as ever,

Salah

Salah A. Soliman

Professor of Pesticide Chemistry & Toxicology

Alexandria University

Senior Expert, Bibliotheca Alexandrina

El-Shatby, Alexandria 21526, EGYPT

E-mail: Salah.Soliman@bibalex.org

Ex. 6 - Personal Privacy

Website: www.bibalex.org

www.bibalex.org/yesbu

Sent from my iPad

Tala[Henry.Tala@epa.gov]; Hanley, Mary[Hanley.Mary@epa.gov] Orme-Zavaleta, Jennifer[Orme-Zavaleta.Jennifer@epa.gov]; Rodan, Bruce[rodan.bruce@epa.gov]; Yamada, Richard (Yujiro)[yamada.richard@epa.gov]; Thayer, Kris[thayer.kris@epa.gov]; Lavoie, Emma[Lavoie.Emma@epa.gov]; Scheifele, Hans[Scheifele.Hans@epa.gov] From: Dourson, Michael Sent: Thur 12/7/2017 8:35:28 PM Subject: RE: Slides we discussed Tina Thanks. Training is important to develop risk assessment erudition. The OPP has an internal senior review board where younger staff can gain their training wheels (so to speak) as their files are peer reviewed. Does NCEA have such a practice? Cheers! Michael... ... L. Dourson, PhD., DABT, FATS, FSRA Senior Advisor to the Administrator U.S. Environmental Protection Agency dourson.michael@epa.gov 202-564-2463 www.epa.gov

Bahadori, Tina[Bahadori.Tina@epa.gov]; Bertrand, Charlotte[Bertrand.Charlotte@epa.gov];

Beck, Nancy[beck.nancy@epa.gov]; Morris, Jeff[Morris.Jeff@epa.gov]; Henry,

To:

From: Bahadori, Tina

Sent: Monday, December 4, 2017 2:59 PM

To: Dourson, Michael <dourson.michael@epa.gov>; Bertrand, Charlotte

<Bertrand.Charlotte@epa.gov>; Beck, Nancy <Beck.Nancy@epa.gov>; Morris, Jeff

<Morris.Jeff@epa.gov>; Henry, Tala <Henry.Tala@epa.gov>; Hanley, Mary

<Hanley.Mary@epa.gov>

Cc: Orme-Zavaleta, Jennifer <Orme-Zavaleta.Jennifer@epa.gov>; Rodan, Bruce

<rodan.bruce@epa.gov>; Yamada, Richard (Yujiro) <yamada.richard@epa.gov>; Thayer, Kris

<thayer.kris@epa.gov>; Lavoie, Emma <Lavoie.Emma@epa.gov>; Scheifele, Hans

<Scheifele.Hans@epa.gov>

Subject: RE: Slides we discussed

Thanks Mike. We have a very strong emphasis on training. Some high level examples include:

- a retreat this week in which our assessment teams will receive hands-on training on evidence synthesis.
- memoranda of understanding with academic institutions, so that students from those institutions can engage in the 'real-world' practice of systematic review and risk assessment as part of their academic training.
- training modules for states and regional risk assessors.

Since Kris joined NCEA, she has made training a very high priority – we have already begun to reap the benefits of her focused investment.

Tina

From: Dourson, Michael

Sent: Monday, December 4, 2017 8:20 AM

To: Bahadori, Tina < Bahadori, Tina@epa.gov >; Bertrand, Charlotte

< Bertrand. Charlotte@epa.gov >; Beck, Nancy < Beck. Nancy@epa.gov >; Morris, Jeff

< Morris.Jeff@epa.gov>; Henry, Tala < Henry.Tala@epa.gov>; Hanley, Mary

< Hanley . Mary @epa.gov >

Cc: Orme-Zavaleta, Jennifer < Orme-Zavaleta.Jennifer@epa.gov >; Rodan, Bruce

<rodan.bruce@epa.gov>; Yamada, Richard (Yujiro) <yamada.richard@epa.gov>; Thayer, Kris

<thayer.kris@epa.gov>; Lavoie, Emma <Lavoie.Emma@epa.gov>; Scheifele, Hans <<u>Scheifele.Hans@epa.gov</u>> Subject: RE: Slides we discussed Tina Thanks for this information. Very helpful. I believe that the OPP has a systematic way of training its younger staff to be better risk assessors. How does NCEA do this please? Cheers! Michael... ... L. Dourson, PhD., DABT, FATS, FSRA Senior Advisor to the Administrator U.S. Environmental Protection Agency dourson.michael@epa.gov 202-564-2463 www.epa.gov From: Bahadori, Tina Sent: Sunday, December 3, 2017 11:23 AM To: Bertrand, Charlotte < Bertrand. Charlotte@epa.gov >; Beck, Nancy < Beck. Nancy@epa.gov >; Dourson, Michael <<u>dourson.michael@epa.gov</u>>; Morris, Jeff <<u>Morris.Jeff@epa.gov</u>>; Henry, Tala < Henry . Tala@epa.gov >; Hanley, Mary < Hanley . Mary@epa.gov > Cc: Orme-Zavaleta, Jennifer < Orme-Zavaleta.Jennifer@epa.gov >; Rodan, Bruce <rodan.bruce@epa.gov>; Yamada, Richard (Yujiro)
yamada.richard@epa.gov>; Thayer, Kris <thayer.kris@epa.gov>; Lavoie, Emma <Lavoie.Emma@epa.gov>; Scheifele, Hans <Scheifele.Hans@epa.gov> Subject: Slides we discussed Dear OCSPP Colleagues, Following up from our 'systematic review' discussions on Friday, I am forwarding the link to the slides we presented at the SAB's Chemical Assessment Advisory Committee (CAAC) meeting in September. We have also presented versions of these materials, in varying detail and depth to other audiences such as in NAS workshops, meeting with the European Food Safety Agency, meetings with state risk assessors, interagency meetings, and scientific conferences. Link to slides: https://yosemite.epa.gov/sab/sabproduct.nsf/AE79F54CBA716293852581A70074264A/\$File/IRIS+Update.pdf We look forward to our continued discussion. Tina Tina Bahadori, Sc.D. Director, National Center for Environmental Assessment (EPA/ORD/NCEA) National Program Director, Human Health Risk Assessment (EPA/ORD/HHRA)

***New RRB Room 71210; Telephone: 202-564-7903; Mobile: 202-680-8771

To: Bowman, Liz[Bowman.Liz@epa.gov]; Jackson, Ryan[jackson.ryan@epa.gov]
Cc: Lyons, Troy[lyons.troy@epa.gov]; Beck, Nancy[beck.nancy@epa.gov]

From: Dourson, Michael

Sent: Thur 10/26/2017 9:59:47 PM
Subject: RE: For Review: Draft Myth v Reality
md2017-10-26 Draft Myth v Reality on Dourson.docx

Here you go!

From: Dourson, Michael

Sent: Thursday, October 26, 2017 5:46 PM

To: Bowman, Liz <Bowman.Liz@epa.gov>; Jackson, Ryan <jackson.ryan@epa.gov>

Cc: Lyons, Troy <lyons.troy@epa.gov>; Beck, Nancy <beck.nancy@epa.gov>

Subject: RE: For Review: Draft Myth v Reality

Dear Colleagues

I will be revising Troy's version momentarily.

Cheers!

Michael

From: Bowman, Liz

Sent: Thursday, October 26, 2017 5:35 PM To: Jackson, Ryan jackson.ryan@epa.gov>

Cc: Lyons, Troy < lyons.troy@epa.gov>; Dourson, Michael < dourson.michael@epa.gov>; Beck,

Nancy < Beck. Nancy@epa.gov>

Subject: RE: For Review: Draft Myth v Reality

Sure

From: Jackson, Ryan

Sent: Thursday, October 26, 2017 4:48 PM **To:** Bowman, Liz <<u>Bowman, Liz@epa.gov</u>>

Cc: Lyons, Troy < lyons.troy@epa.gov>; Dourson, Michael < dourson.michael@epa.gov>; Beck,

Nancy < Beck. Nancy@epa.gov>

Subject: Re: For Review: Draft Myth v Reality

Do you want to get together on this this afternoon?

Ryan Jackson

Chief of Staff

U.S. EPA

Ex. 6 - Personal Privacy

On Oct 26, 2017, at 3:35 PM, Bowman, Liz < Bowman.Liz@epa.gov > wrote:

Michael, did you send me edits? You mentioned you did, but I don't have an email from you?

From: Lyons, Troy

Sent: Thursday, October 26, 2017 3:18 PM

To: Bowman, Liz < Bowman.Liz@epa.gov >; Jackson, Ryan < jackson.ryan@epa.gov >; Dourson, Michael < dourson.michael@epa.gov >; Beck, Nancy < Beck.Nancy@epa.gov >

Subject: RE: For Review: Draft Myth v Reality

Take a look at my edits. I thought it would be good to highlight the credentials upfront and then again in the MvF

From: Bowman, Liz

Sent: Thursday, October 26, 2017 1:54 PM

To: Jackson, Ryan < jackson.ryan@epa.gov>; Dourson, Michael

<<u>dourson.michael@epa.gov</u>>; Lyons, Troy <<u>lyons.troy@epa.gov</u>>; Beck, Nancy <<u>Beck.Nancy@epa.gov</u>>

Subject: For Review: Draft Myth v Reality

Attached.

Liz Bowman

U.S. Environmental Protection Agency (EPA)

Office: 202-564-3293

From: Dourson, Michael Sent: Thur 11/16/2017 1:10:02 AM Subject: Re: Chlopyrifos Kaitlin No, let's try electronic. Michael Sent from my iPad On Nov 15, 2017, at 11:30 AM, Keller, Kaitlin < keller.kaitlin@epa.gov> wrote: Quick question—do you want hard copies of the SAP reports and studies in addition to the electronic copies sent yesterday? Thanks, Kaitlin From: Keller, Kaitlin Sent: Tuesday, November 14, 2017 5:49 PM To: Beck, Nancy <beck.nancy@epa.gov> Cc: Dourson, Michael <<u>dourson.michael@epa.gov</u>>; Bertrand, Charlotte <Bertrand.Charlotte@epa.gov> Subject: RE: Chlopyrifos Nancy, Attached are the last three Chlorpyrifos SAP reports. Also attached is the 2016 OPP systematic review and an appendix document from the 2012 SAP that I thought looked like a good high level snapshot of the different data.

I'm also about to send 4 additional emails with Ex. 5 - Deliberative Process

Ex. 5 - Deliberative Process

To:

Keller, Kaitlin[keller.kaitlin@epa.gov]

Ex. 5 - Deliberative Process vill have something I'm still working on before the Friday briefing though. Charlotte & Mike—Let me know if you'd like hard copies and we'll get those to you tomorrow. Thanks, Kaitlin From: Beck, Nancy Sent: Friday, November 10, 2017 2:20 PM To: Keller, Kaitlin < keller.kaitlin@epa.gov> Cc: Dourson, Michael dourson, Michael dourson.michael@epa.gov>; Bertrand, Charlotte <Bertrand.Charlotte@epa.gov> Subject: Chlopyrifos Kaitlyn, Ex. 5 - Deliberative Process Before next weeks briefing can you send us Ex. 5 - Deliberative Process vould Ex. 5 - Deliberative Process I think I have all this but my computer access is limited. I won't need printouts as I have them already. Ex. 5 - Deliberative Process Also, if you have time

Ex. 5 - Deliberative Process

Thanks.			
Sent from my iPhone, please ex	xcuse typos.		

To: Weiss, Steven[Weiss.Steven@epa.gov]; Bertrand, Charlotte[Bertrand.Charlotte@epa.gov] Jewell, Shannon[jewell.shannon@epa.gov]; Sinclair, Geoffrey[Sinclair.Geoffrey@epa.gov]; Cc: Keigwin, Richard[Keigwin.Richard@epa.gov]; Isbell, Diane[Isbell.Diane@epa.gov]; Parsons, Laura[Parsons.Laura@epa.gov]; Kyprianou, Rose[Kyprianou.Rose@epa.gov]; Hebert, John[Hebert.John@epa.gov]; Mitchell, Emily[Mitchell.Emily@epa.gov]; Keller, Kaitlin[keller.kaitlin@epa.gov] From: Dourson, Michael Thur 11/30/2017 1:09:25 PM Sent: Subject: RE: AD Division 101 w/Charlotte Bertrand & Mike Dourson Steve Thanks. Looking forward to the briefing. Cheers! Michael... ... L. Dourson, PhD., DABT, FATS, FSRA Senior Advisor to the Administrator U.S. Environmental Protection Agency dourson.michael@epa.gov 202-564-2463 www.epa.gov

From: Weiss, Steven

Sent: Wednesday, November 29, 2017 6:02 PM

To: Bertrand, Charlotte <Bertrand.Charlotte@epa.gov>; Dourson, Michael

<dourson.michael@epa.gov>

Cc: Jewell, Shannon <jewell.shannon@epa.gov>; Sinclair, Geoffrey

<Sinclair.Geoffrey@epa.gov>; Keigwin, Richard <Keigwin.Richard@epa.gov>; Isbell, Diane

<Isbell.Diane@epa.gov>; Parsons, Laura <Parsons.Laura@epa.gov>; Kyprianou, Rose

<Kyprianou.Rose@epa.gov>; Hebert, John <Hebert.John@epa.gov>; Mitchell, Emily

<Mitchell.Emily@epa.gov>; Keller, Kaitlin <keller.kaitlin@epa.gov>

Subject: RE: AD Division 101 w/Charlotte Bertrand & Mike Dourson

Hi Charlotte and Mike,

Attached are the slides for tomorrow's mtg.

Thanks,

Steve

Steven H. Weiss
Acting Director
Antimicrobials Division
Office of Pesticide Programs
US Environmental Protection Agency
Weiss.Steven@epa.gov
(703)308-8293

----Original Appointment-----**From:** Bertrand, Charlotte

Sent: Tuesday, November 07, 2017 2:39 PM

To: Bertrand, Charlotte; Weiss, Steven; Hebert, John; Kyprianou, Rose; Mitchell, Emily;

Parsons, Laura; Dourson, Michael

Cc: Jewell, Shannon; Sinclair, Geoffrey; Keigwin, Richard; Isbell, Diane

Subject: AD Division 101 w/Charlotte Bertrand & Mike Dourson

When: Thursday, November 30, 2017 11:00 AM-11:30 AM (UTC-05:00) Eastern Time (US &

Canada).

Where: RM 12621 PY South

To: From: Sent: Subject:	Washington, Valerie[Washington.Valerie@epa.gov] Dourson, Michael Tue 10/31/2017 2:39:58 PM Re: Training
Valerie	
the 5 pm.	eetings all day. Do you want my calendar? I should be back in the office today around Tomorrow is not much better. Perhaps we can work on this first thing tomorrow or sometime Thursday afternoon.
Cheers!	
Michael	
Sent from	n my iPad
On Oct 3	1, 2017, at 10:19 AM, Washington, Valerie < Washington. Valerie@epa.gov > wrote:
Good I	Morning Michael,
	know when you are coming back to the office we need to register you for your g but you have to go online and register.
Sorry	
Thanks	\mathbf{S}

To: Anderson, Brian[Anderson.Brian@epa.gov]; Nesci, Kimberly[Nesci.Kimberly@epa.gov];

Bertrand, Charlotte[Bertrand.Charlotte@epa.gov]; Keller, Kaitlin[keller.kaitlin@epa.gov]

Cc: Keigwin, Richard[Keigwin.Richard@epa.gov]; Echeverria,

Marietta[Echeverria.Marietta@epa.gov]
From: Dourson, Michael

Sent: Thur 12/7/2017 8:31:03 PM
Subject: RE: Follow-up from EFED 101

Brian

Thanks. This is very helpful.

Cheers!

Michael...

... L. Dourson, PhD., DABT, FATS, FSRA

Senior Advisor to the Administrator

U.S. Environmental Protection Agency

dourson.michael@epa.gov

202-564-2463

www.epa.gov

From: Anderson, Brian

Sent: Tuesday, December 5, 2017 4:43 PM

To: Dourson, Michael <dourson.michael@epa.gov>; Nesci, Kimberly <Nesci.Kimberly@epa.gov>; Bertrand, Charlotte <Bertrand.Charlotte@epa.gov>; Keller, Kaitlin <keller.kaitlin@epa.gov>
Cc: Keigwin, Richard <Keigwin.Richard@epa.gov>; Echeverria, Marietta <Echeverria.Marietta@epa.gov>
Subject: Follow-up from EFED 101

Hi Michael,

Ex. 5 - Deliberative Process

Please let me know if you need any more information or have any additional questions.

Thanks,

Brian

To: Morris, Jeff[Morris.Jeff@epa.gov]

From: Dourson, Michael

Sent: Tue 11/21/2017 6:10:53 PM

Subject: RE: SNURs and Risk Evaluation/PBT Scoping (Call in Ex. 6 - Personal Privacy

Jeff

Ex. 5 - Deliberative Process

Cheers!

Michael

From: Morris, Jeff

Sent: Tuesday, November 21, 2017 12:55 PM To: Dourson, Michael <dourson.michael@epa.gov>

Subject: RE: SNURs and Risk Evaluation/PBT Scoping (Call in Ex. 6 - Personal Privacy

Ex. 6 - Personal Privacy

Michael,

Yes, should be a good discussion. One thing I'm going to raise at the meeting is whether, for the

Ex. 5 - Deliberative Process

SNURs are not risk based.

Ex. 5 - Deliberative Process

Ex. 5 - Deliberative Process

Jeff

From: Dourson, Michael

Sent: Tuesday, November 21, 2017 12:38 PM To: Morris, Jeff < Morris. Jeff@epa.gov>

Cc: Anderson, Monique <anderson.monique@epa.gov>; Vendinello, Lynn

- <Vendinello.Lynn@epa.gov>; Wolf, Joel <Wolf.Joel@epa.gov>; Anderson, Steve
- <Anderson.Steve@epa.gov>; Frazer, Brian <Frazer.Brian@epa.gov>; Baptist, Erik
- <<u>baptist.erik@epa.gov</u>>; Mclean, Kevin <<u>Mclean, Kevin@epa.gov</u>>; Grant, Brian
- <Grant.Brian@epa.gov>; Wills, Jennifer < Wills.Jennifer@epa.gov>; Fisher, Bethany
- <fisher.bethany@epa.gov>; Aranda, Amber <aranda.amber@epa.gov>; Beck, Nancy
- < <u>Beck.Nancy@epa.gov</u>>; Bertrand, Charlotte < <u>Bertrand.Charlotte@epa.gov</u>>; Mottley, Tanya
- <Mottley.Tanya@epa.gov>; Doa, Maria <Doa.Maria@epa.gov>; Henry, Tala
- <Henry.Tala@epa.gov>

Subject: RE: SNURs and Risk Evaluation/PBT Scoping (Call in Ex. 6 - Personal Privacy Ex. 6 - Personal Privacy

Jeff

I am looking forward to this discussion, but have a few big-picture questions that would help me

Ex. 5 - Deliberative Process

Thanks for considering these questions prior to the briefing and for me taking so long to get up to speed.

Cheers!

Michael...

... L. Dourson, PhD., DABT, FATS, FSRA

Senior Advisor to the Administrator

U.S. Environmental Protection Agency

dourson.michael@epa.gov

202-564-2463

www.epa.gov

----Original Appointment----

From: Motley, Judy On Behalf Of Baptist, Erik Sent: Tuesday, November 21, 2017 10:32 AM

To: Mclean, Kevin; Grant, Brian; Wills, Jennifer; Fisher, Bethany; Aranda, Amber; Beck,

Nancy; Bertrand, Charlotte; Morris, Jeff; Mottley, Tanya; Doa, Maria; Henry, Tala

Cc: Anderson, Monique; Vendinello, Lynn; Wolf, Joel; Dourson, Michael; Anderson, Steve;

Frazer, Brian

Subject: SNURs and Risk Evaluation/PBT Scoping (Call in Ex. 6 - Personal Privacy

When: Tuesday, November 21, 2017 1:00 PM-2:00 PM (UTC-05:00) Eastern Time (US &

Canada).

Where: DCRoomARN4045/DC-Ariel-Rios-OGC

Call in: Ex. 6 - Personal Privacy

To: Krasnic, Toni[krasnic.toni@epa.gov]

Cc: Beck, Nancy[Beck.Nancy@epa.gov]; Bertrand, Charlotte[Bertrand.Charlotte@epa.gov]; Morris,

Jeff[Morris.Jeff@epa.gov]; Doa, Maria[Doa.Maria@epa.gov]; Wolf, Joel[Wolf.Joel@epa.gov]

From: Dourson, Michael

Sent: Fri 11/17/2017 11:29:53 PM

Subject: Re: TCE

Toni

Thanks!

Michael

Sent from my iPhone

On Nov 17, 2017, at 12:57 PM, Krasnic, Toni < krasnic.toni@epa.gov > wrote:

Hi Michael,

Thank you for your interest in this topic. We'll pull together additional information on this visit and will send it to you once it's compiled.

Thanks,

Toni Krasnic

Existing Chemicals Branch

EPA/OCSPP/OPPT/CCD/ECB

WJC East, 4134D | (202) 564-0984

From: Dourson, Michael

Sent: Friday, November 17, 2017 12:37 PM **To:** Krasnic, Toni <<u>krasnic.toni@epa.gov</u>>

Cc: Beck, Nancy <<u>Beck.Nancy@epa.gov</u>>; Bertrand, Charlotte <<u>Bertrand.Charlotte@epa.gov</u>>; Morris, Jeff <<u>Morris.Jeff@epa.gov</u>>

Subject: TCE Dear Toni Krasnic I would be interested in more information regarding this topic. Visit at Integer facility on use of TCE in vapor degreasing in the manufacture of medical devices: On October 19, 2017, CCD, CESSD, and RAD staff visited Integer's facility in Minneapolis, MN. Integer was formed from the merger of Greatbatch, Lake Region Medical, and Electrochem, and specializes in the design and development of medical devices and power solutions for the medical and non-medical markets. EPA toured Integer's facility where they make various medical devices, for a demonstration of open-top vapor degreasing, spray degreasing, enclosed vapor degreasing, and aqueous degreasing. Integer also provided a tour of their manufacturing process of medical devices and showed samples of their actual medical products. Additionally, they discussed the research and testing they've done on non-TCE alternatives and Ex. 5 - Deliberative Process Ex. 5 - Deliberative Process Thanks! Michael... ... L. Dourson, PhD., DABT, FATS, FSRA Senior Advisor to the Administrator U.S. Environmental Protection Agency dourson.michael@epa.gov 202-564-2463

www.epa.gov

Cc: Beck, Nancy[Beck.Nancy@epa.gov]
To: Morris, Jeff[Morris.Jeff@epa.gov]

From: Dourson, Michael

Sent: Tue 10/24/2017 5:27:15 PM

Subject: Re: PRE-PRIORIZATION PUBLIC MEETING

Jeff

I am currently scheduled for a session at the Society for Risk Analysis meeting from 8:30 to 10 am on 12/11. This commitment was made well before my nomination for AA. What did you have in mind for my activity at your meeting, if anything?

Cheers!

Michael

Sent from my iPad

> On Oct 24, 2017, at 1:00 PM, Morris, Jeff < Morris.Jeff@epa.gov> wrote:

>

- > Additional information will be provided.
- > <meeting.ics>

To: Thayer, Kris[thayer.kris@epa.gov]; Beck, Nancy[beck.nancy@epa.gov]; Yamada, Richard (Yujiro)[yamada.richard@epa.gov]; Rodan, Bruce[rodan.bruce@epa.gov]; Bahadori, Tina[Bahadori.Tina@epa.gov]; Morris, Jeff[Morris.Jeff@epa.gov]; Henry, Tala[Henry.Tala@epa.gov]; Orme-Zavaleta, Jennifer[Orme-Zavaleta.Jennifer@epa.gov]; Hanley, Mary[Hanley.Mary@epa.gov]; Bertrand, Charlotte[Bertrand.Charlotte@epa.gov]; Scheifele, Hans[Scheifele.Hans@epa.gov]; Lavoie, Emma[Lavoie.Emma@epa.gov]; Camacho, Iris[Camacho.Iris@epa.gov]; Bateson, Thomas[Bateson.Thomas@epa.gov]; Kraft, Andrew[Kraft.Andrew@epa.gov]

From: Dourson, Michael

Sent: Wed 12/20/2017 1:45:18 PM

Subject: Systematic Review and Risk Assessment

Kris and Colleagues

I very much appreciate the meeting yesterday. **Ex. 5 - Deliberative Process**

Ex. 5 - Deliberative Process

Ex. 5 - Deliberative Process

Cheers!

Michael...

... L. Dourson, PhD., DABT, FATS, FSRA

Senior Advisor to the Administrator

U.S. Environmental Protection Agency

dourson.michael@epa.gov

202-564-2463

www.epa.gov

From: Thayer, Kris

Sent: Wednesday, December 20, 2017 3:36 AM

To: Beck, Nancy <Beck.Nancy@epa.gov>; Yamada, Richard (Yujiro)

<yamada.richard@epa.gov>; Rodan, Bruce <rodan.bruce@epa.gov>; Bahadori, Tina

<Bahadori.Tina@epa.gov>; Dourson, Michael <dourson.michael@epa.gov>; Morris, Jeff

<Morris.Jeff@epa.gov>; Henry, Tala <Henry.Tala@epa.gov>; Orme-Zavaleta, Jennifer <Orme-</p>

Zavaleta.Jennifer@epa.gov>; Hanley, Mary <Hanley.Mary@epa.gov>; Bertrand, Charlotte

<Bertrand.Charlotte@epa.gov>; Scheifele, Hans <Scheifele.Hans@epa.gov>; Lavoie, Emma

<Lavoie.Emma@epa.gov>; Camacho, Iris <Camacho.Iris@epa.gov>; Bateson, Thomas

<Bateson.Thomas@epa.gov>; Kraft, Andrew < Kraft.Andrew@epa.gov>

Subject: Background information on study scoring in systematic review

Thanks again for the meeting today. Hopefully both of our programs will benefit from the robust discussion.

As noted today, I am headed out of the country so will unlikely be able to make another meeting this week. However, I wanted to quickly share some of the systematic review guidance materials

Ex. 5 - Deliberative Process

2011 Institute of Medicine report (where the TSCA definition of systematic review is taken

Ex. 5 - Deliberative Process

Cochrane Handbook (the most recognized source of systematic review guidance)

http://handbook-5-1.cochrane.org/chapter 8/8 3 3 quality scales and cochrane reviews.htm

Ex. 5 - Deliberative Process

Ex. 5 - Deliberative Process

Ex. 5 - Deliberative Process

Ex. 5 - Deliberative Process

Happy holidays! -k

Kristina Thayer, Ph.D.

Director, Integrated Risk Information System (IRIS) Division

National Center for Environmental Assessment, NCEA

ORD, USEPA

Mail Code: B243-01

Building: Bldg B (Room B211I)

Research Triangle Park, NC 27711

(919) 541-0152 RTP

(202) 564-1771 Ronald Reagan Building (room 51203)

Skype: kristina.thayer

thayer.kris@epa.gov

Cc: Beck, Nancy[Beck.Nancy@epa.gov]
To: Jackson, Ryan[jackson.ryan@epa.gov]

From: Dourson, Michael

Sent: Mon 10/23/2017 10:11:14 AM

Subject: Flight delay

Ryan

My flight out of Ohio has been delayed, which will cause me to be late for your meeting.

Sorry!

Michael

Sent from my iPad

To: From: Sent: Subject:	Beck, Nancy[beck.nancy@epa.gov] Dourson, Michael Mon 12/18/2017 6:08:37 PM RE: TRI Analysis Release Briefing
Thanks, I	will be there.
Sent: Mo To: Dour Cc: Berti	eck, Nancy onday, December 18, 2017 1:06 PM rson, Michael <dourson.michael@epa.gov> rand, Charlotte <bertrand.charlotte@epa.gov> Re: TRI Analysis Release Briefing</bertrand.charlotte@epa.gov></dourson.michael@epa.gov>
It's an intattend.	formational briefing for OPA and the office of engagement. You are welcome to
Deputy A P: 202-56 M: Ex.6-Pe	Beck, Ph.D., DABT assistant Administrator, OCSPP 64-1273 arsonal Privacy ncy@epa.gov
On Dec 1	8, 2017, at 1:02 PM, Dourson, Michael < dourson.michael@epa.gov > wrote:
Nancy	
Please	let me know if you need me at this briefing.
Cheers	!
Mike	
Or	iginal Appointment

From: Marshall, Venus On Behalf Of Beck, Nancy

Sent: Monday, December 18, 2017 7:25 AM

To: Bowman, Liz; Block, Molly; Hewitt, James; Bennett, Tate; Briere, Caitlin; Reisman, Larry; Devito, Steve; Turk, David; Tomassoni, Guy; Swenson, Sarah; Berckes, Nicole; Hartman, Mark; Bertrand, Charlotte; Dourson, Michael

Cc: Strauss, Linda; Ortiz, Julia; Pierce, Alison; Blunck, Christopher; Bolen, Derrick

Subject: TRI Analysis Release Briefing

When: Monday, December 18, 2017 3:00 PM-4:00 PM (UTC-05:00) Eastern Time (US & Canada).

Where: DCRoomEast 3371/DC-EPA-EAST-OCSPP (Call in Ex. 6 - Personal Privacy , access code

Ex. 6 - Personal Privacy

Bowman, Liz[Bowman.Liz@epa.gov]; Jackson, Ryan[jackson.ryan@epa.gov]; Lyons, To: Troy[lyons.troy@epa.gov]; Beck, Nancy[beck.nancy@epa.gov] Dourson, Michael From: Thur 10/26/2017 9:44:46 PM Sent: Subject: RE: For Review: Draft Myth v Reality md2017-10-26 Draft Myth v Reality on Dourson.docx Liz Very nice. Please see the attached tweaked version (sorry, but the strike and replace version did not carry over). Cheers! Michael From: Bowman, Liz Sent: Thursday, October 26, 2017 1:54 PM To: Jackson, Ryan <jackson.ryan@epa.gov>; Dourson, Michael <dourson.michael@epa.gov>; Lyons, Troy ! Beck, Nancy Beck, Nancy <a href="mailto: **Subject:** For Review: Draft Myth v Reality

Attached.

Liz Bowman

U.S. Environmental Protection Agency (EPA)

Office: 202-564-3293

To: Beck, Nancy[beck.nancy@epa.gov]

From: Dourson, Michael

Sent: Wed 11/1/2017 10:10:07 PM

Subject: RE: Do I

Thanks!

----Original Message-----From: Beck, Nancy

Sent: Wednesday, November 1, 2017 3:02 PM To: Dourson, Michael dourson.michael@epa.gov

Subject: RE: Do I

Bernhardt

Nancy B. Beck, Ph.D., DABT
Deputy Assistant Administrator, OCSPP
P: 202-564-1273

M: Ex. 6 - Personal Privacy beck.nancy@epa.gov

----Original Message----From: Dourson, Michael

Sent: Wednesday, November 1, 2017 2:50 PM To: Beck, Nancy <Beck.Nancy@epa.gov>

Subject: Do I

Please call asap. Need doi contact.

Mike

Sent from my iPad

To: Hanley, Mary[Hanley.Mary@epa.gov]; Morris, Jeff[Morris.Jeff@epa.gov]; Bertrand,

Charlotte[Bertrand.Charlotte@epa.gov]; Wise, Louise[Wise.Louise@epa.gov]

Cc: Beck, Nancy[beck.nancy@epa.gov]

From: Dourson, Michael

Sent: Tue 11/7/2017 6:21:47 PM

Subject: RE: Paragraphs

Mary

Very nice.

Michael

----Original Message-----From: Hanley, Mary

Sent: Tuesday, November 7, 2017 1:14 PM

To: Morris, Jeff , Bertrand, Charlotte , Dourson, Jeff , Dourson, Jeff <a href="Morris.Jeff"

Michael <dourson.michael@epa.gov>; Wise, Louise <Wise.Louise@epa.gov>

Cc: Beck, Nancy <Beck.Nancy@epa.gov>

Subject: RE: Paragraphs

Let me know if this works for this topic. Charlotte I know this is longer than a few short bullets...

Ex. 5 - Deliberative Process

----Original Message----

From: Morris, Jeff

Sent: Tuesday, November 07, 2017 12:31 PM

To: Bertrand, Charlotte <Bertrand.Charlotte@epa.gov>; Dourson, Michael <dourson.michael@epa.gov>;

Wise, Louise <Wise.Louise@epa.gov>; Hanley, Mary <Hanley.Mary@epa.gov>

Cc: Beck, Nancy <Beck.Nancy@epa.gov>

Subject: RE: Paragraphs

Ex. 5 - Deliberative Process

Ex. 5 - Deliberative Process

----Original Message----

From: Bertrand, Charlotte

Sent: Tuesday, November 07, 2017 11:52 AM

To: Dourson, Michael <dourson.michael@epa.gov>; Morris, Jeff <Morris.Jeff@epa.gov>; Wise, Louise

<Wise.Louise@epa.gov>; Hanley, Mary <Hanley.Mary@epa.gov>

Cc: Beck, Nancy <Beck.Nancy@epa.gov>

Subject: RE: Paragraphs

Thank you. I spoke to RJ, he's asked for fees and legacy uses as well. Mary is working on this. Need to be done by mid-afternoon.

----Original Message----

From: Dourson, Michael

Sent: Tuesday, November 07, 2017 9:59 AM

To: Bertrand, Charlotte <Bertrand.Charlotte@epa.gov>; Morris, Jeff <Morris.Jeff@epa.gov>; Wise,

Louise <Wise.Louise@epa.gov>

Subject: Paragraphs

Jeff, Charlotte and Louise

Ex. 5 - Deliberative Process

Cheers!

Michael

Sent from my iPad

To: From: Sent: Subject:	Beck, Nancy[beck.nancy@epa.gov] Dourson, Michael Mon 12/18/2017 6:08:11 PM RE: IRIS Agency Review - Ex.5-Deliberative Process		
Ok, but let's see if they allow me access first.			
Sent: Mo	eck, Nancy onday, December 18, 2017 1:05 PM rson, Michael <dourson.michael@epa.gov> Re: IRIS Agency Review - Ex.5-Deliberative Process</dourson.michael@epa.gov>		
I can lend	d you my hard copy if that's easier.		
Deputy A P: 202-56 M: Ex.6-Pe	Beck, Ph.D., DABT Assistant Administrator, OCSPP 64-1273 Personal Privacy hcy@epa.gov		
On Dec 1	8, 2017, at 1:00 PM, Dourson, Michael < <u>dourson.michael@epa.gov</u> > wrote:		
Nancy			
Thanks	s for sending me this link. I requested access.		
Cheers	ş!		
Mike			
Sent: 1	Beck, Nancy Monday, December 18, 2017 12:11 PM ourson, Michael < <u>dourson.michael@epa.gov</u> >		

Subject: FW: IRIS Agency Review - Ex. 5 - Deliberative Process

Nancy B. Beck, Ph.D., DABT

Deputy Assistant Administrator, OCSPP

P: 202-564-1273

M: Ex. 6 - Personal Privacy

beck.nancy@epa.gov

From: Soto, Vicki

Sent: Friday, December 8, 2017 2:26 PM

To: Miller, Gregory < Miller. Gregory@epa.gov >; Lowit, Anna < Lowit. Anna@epa.gov >; Laessig, Susan < Laessig. Susan@epa.gov >; Hamernik, Karen < Hamernik. Karen@epa.gov >;

Raffaele, Kathleen < raffaele.kathleen@epa.gov >; Murphy, Deirdre

< <u>Murphy.Deirdre@epa.gov</u>>; Hoyer, Marion < <u>hoyer.marion@epa.gov</u>>; Vasu, Amy

< Vasu. Amy@epa.gov>; Axelrad, Daniel < Axelrad. Daniel@epa.gov>; Dzubow, Rebecca

<<u>Dzubow.Rebecca@epa.gov</u>>; Cassidy, Meghan <<u>Cassidy.Meghan@epa.gov</u>>; Olsen,

Marian < Olsen.Marian@epa.gov>; Gehlhaus, Martin < Gehlhaus.Martin@epa.gov>;

Pollard, Solomon < Pollard. Solomon@epa.gov >; Adams, Glenn < Adams. Glenn@epa.gov >;

Mangino, Mario < mangino.mario@epa.gov >; Milburn, Anna < Milburn.Anna@epa.gov >; Schumacher Kelly < Schumac

Schumacher, Kelly < Schumacher.Kelly@epa.gov >; Griffin, Susan

< <u>Griffin.Susan@epa.gov</u>>; Hiatt, Gerald < <u>Hiatt.Gerald@epa.gov</u>>; Kissinger, Lon

< Kissinger.Lon@epa.gov>; Axelrad, Daniel < Axelrad.Daniel@epa.gov>; Barone, Stan

< <u>Barone.Stan@epa.gov</u>>; Schappelle, Seema < <u>Schappelle.Seema@epa.gov</u>>; Markey,

Kristan < Markey.Kristan@epa.gov>; Camacho, Iris < Camacho, Iris@epa.gov>; Henry, Tala

< Henry. Tala@epa.gov >; Lowit, Anna < Lowit. Anna@epa.gov >; Foster, Stiven

<<u>Foster.Stiven@epa.gov</u>>; Raffaele, Kathleen <<u>raffaele.kathleen@epa.gov</u>>; Murphy,

Deirdre < Murphy. Deirdre@epa.gov >; Mazza, Carl < Mazza. Carl@epa.gov >; Ohanian,

Edward < Ohanian. Edward@epa.gov >; Strong, Jamie < Strong. Jamie@epa.gov >; Firestone,

Michael <<u>Firestone.Michael@epa.gov</u>>; Gibbons, Catherine

<Gibbons.Catherine@epa.gov>; Radke-Farabaugh, Elizabeth <radke-</p>

farabaugh.elizabeth@epa.gov>; Kraft, Andrew < Kraft.Andrew@epa.gov>; Cogliano,

Vincent <cogliano.vincent@epa.gov>; Sams, Reeder <Sams.Reeder@epa.gov>; Birchfield,

```
Norman <Birchfield.Norman@epa.gov>; Glenn, Barbara <Glenn.Barbara@epa.gov>;
Hogan, Karen < Hogan. Karen@epa.gov>; Pratt, Margaret < pratt.margaret@epa.gov>; Luke,
April <<u>Luke.April@epa.gov</u>>; Woodall, George <<u>Woodall.George@epa.gov</u>>;
'Patel.Molini@epa.gov' <Patel.Molini@epa.gov>; Flowers, Lynn
<<u>Flowers.Lynn@epa.gov</u>>; Fritz, Jason <<u>Fritz.Jason@epa.gov</u>>; Olsen, Marian
<Olsen.Marian@epa.gov>; Rogers, John M. <Rogers.John@epa.gov>; Vasu, Amy
<Vasu.Amy@epa.gov>; Dzubow, Rebecca <Dzubow.Rebecca@epa.gov>; Newhouse,
Kathleen <Newhouse.Kathleen@epa.gov>; Kirk, Andrea <Kirk.Andrea@epa.gov>;
Hospital, Jocelyn < Hospital. Jocelyn@epa.gov>; Dishaw, Laura < Dishaw. Laura@epa.gov>;
Galizia, Audrey < Galizia. Audrey@epa.gov >; Persad, Amanda
<Persad.Amanda@epa.gov>; Dockins, Chris <Dockins.Chris@epa.gov>; Hotchkiss,
Andrew < Hotchkiss Andrew@epa.gov >; Fallace, Katherine < fallace.katherine@epa.gov >;
Yaquian-Luna, Jose < yaquian-luna.josea@epa.gov >; Druwe, Ingrid
<Druwe.Ingrid@epa.gov>; Davis, Allen <Davis.Allen@epa.gov>; Beck, Nancy
<<u>Beck.Nancy@epa.gov</u>>; Pardo, Larissa <<u>Pardo.Larissa@epa.gov</u>>; Sasso, Alan
<Sasso.Alan@epa.gov>; Berner, Ted <Berner.Ted@epa.gov>; Stohs, Sheryl
<stohs.sheryl@epa.gov>; Griffiths, Charles <Griffiths.Charles@epa.gov>; Hodes, Colette
< Hodes. Colette@epa.gov>; Morris, Jeff < Morris. Jeff@epa.gov>; Soares, Barbara
<soares.barbara@epa.gov>; Congleton, Johanna <congleton.johanna@epa.gov>; Owens,
Beth < Owens. Beth@epa.gov>
Cc: Burgess, Michele < Burgess. Michele@epa.gov >; Euling, Susan
<Euling.Susan@epa.gov>; Bussard, David <Bussard.David@epa.gov>; Thayer, Kris
<thayer.kris@epa.gov>; Ramasamy, Santhini < Ramasamy.Santhini@epa.gov>; Avery,
James <Avery James@epa.gov>; Shams, Dahnish <Shams.Dahnish@epa.gov>; Soto, Vicki
<Soto.Vicki@epa.gov>; Lavoie, Emma <Lavoie.Emma@epa.gov>; D'Amico, Louis
<DAmico.Louis@epa.gov>; Bahadori, Tina <Bahadori.Tina@epa.gov>; Vandenberg, John
<Vandenberg.John@epa.gov>; Jones, Samantha <Jones.Samantha@epa.gov>; Gatchett,
Annette <Gatchett. Annette@epa.gov>; Rieth, Susan <Rieth.Susan@epa.gov>; Morozov,
Viktor < Morozov. Viktor@epa.gov >; Subramaniam, Ravi < Subramaniam. Ravi@epa.gov >;
Lee, Janice < Lee. Janice S@epa.gov >; Hawkins, Belinda < Hawkins. Belinda@epa.gov >;
Radke-Farabaugh, Elizabeth < radke-farabaugh.elizabeth@epa.gov>
Subject: IRIS Agency Review - Ex. 5 - Deliberative Process
```

Dear IRIS Agency Reviewers –

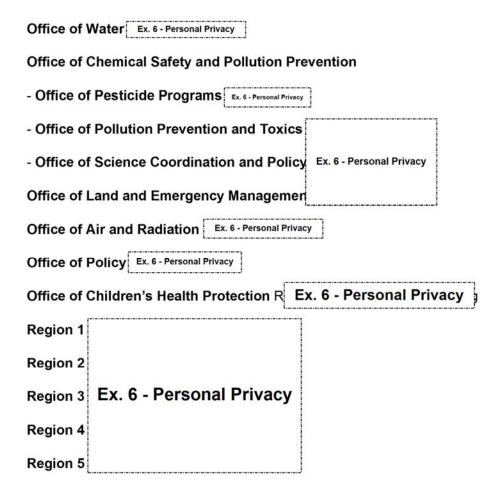
We are pleased to provide our Ex. 5 - Deliberative Process for your review. This document is being shared with the group that has been meeting with the IRIS Program to discuss systematic review, and so is broader than the usual review group.

We are now sharing all review documents on the IRIS Agency SharePoint site, located here:

https://usepa.sharepoint.com/sites/ORD_Work/IRISagencyreview/SitePages/Home.aspx (see the Review Documents section). Three specific notes for this review are provided below:

- Please focus comments only on substantive issues.
- Combine all comments from an office or region into a single document (see Agency Review contact list below), then upload to the Systematic Review folder on the SharePoint site.
- Post all comments to the SharePoint site by <u>COB Friday</u>, <u>January 12</u>

Thank you for your participation in EPA's IRIS Program. We appreciate your scientific input on IRIS draft documents. Please let us know if you have any questions.



Region 6	
Region 7	
Region 8	Ex. 6 - Personal Privacy
Region 9	
Region 1	1

Vicki Soto | 202-564-3077 (new!) | soto.vicki@epa.gov ORD/NCEA

Mailing Address:

 $\begin{tabular}{l} USEPA Headquarters | William Jefferson Clinton Building | 1200 Pennsylvania Avenue, N.W. | MC: 8601P | Washington, DC 20460 \\ \end{tabular}$

To: Bowman, Liz[Bowman.Liz@epa.gov]

From: Dourson, Michael

Sent: Thur 10/26/2017 8:42:30 PM **Subject:** Fwd: Collaborative Works

Excerpts of TERA's Collaborative Work 9-28-17.docx

ATT00001.htm

Liz

Here it is again. Sorry!

Michael

Sent from my iPhone

Begin forwarded message:

From: "Dourson, Michael" < dourson.michael@epa.gov>

Date: October 26, 2017 at 11:58:03 AM EDT **To:** "Bowman, Liz" < Bowman, Liz@epa.gov>

Cc: "Lyons, Troy" < lyons.troy@epa.gov >, "Beck, Nancy" < Beck.Nancy@epa.gov >,

"Jackson, Ryan" <<u>jackson.ryan@epa.gov</u>> **Subject: Fwd: Collaborative Works**

Liz

Here is an extended version of the summary response I did for Carper's table that gives additional details for his 8 chemicals and other issues that were raised in various websites. These additional topics may or may not be helpful.

I would be happy to expand on any of these topics, or to respond to additional issues as needed.

Cheers!

Michael

Sent from my iPad

Begin forwarded message:

From: "Maier, Michael (maierma)" < maierma@ucmail.uc.edu>

Date: October 26, 2017 at 11:46:08 AM EDT

To: "dourson.michael@epa.gov" <dourson.michael@epa.gov>

Subject: Collaborative Works

Example of Collaborative Work in Environmental Risk Assessment by Toxicology Excellence for Risk Assessment (TERA) and the Risk Science Center of the University of Cincinnati, College of Medicine

TERA was founded on the belief that an independent non-profit organization can provide a unique function to protect human health by conducting scientific research and development on risk issues in a transparent and collaborative fashion. One-third of TERA/RSC effort has been for industries; 2/3 has been for government groups. The projects below are examples of this transparent and/or collaborative work.

CPSC: Draft Final Rule ¹

Toxicology Excellence for Risk Assessment (TERA) and the Risk Science Center of the University of Cincinnati, College of Medicine are contractors to the Consumer Products Safety Commission. Below is a recent public exchange that might warrant the EPW committee attention.

<u>Public Comment 16</u>: A commenter states that the contractor (TERA) engaged by the CPSC to study phthalate use and investigate the presence of phthalates in four specified plastics may have a conflict of interest. The commenter notes TERA's past litigation support for regulated industries. The commenter asserts TERA's potential conflict of interest is exemplified in a 2016 paper sponsored by a chemical manufacturers' trade group.

The commenter adds that TERA is a founding member of the Alliance for Risk Assessment (ARA). The ARA's Standing Panel includes the TERA founder, two industry consultants, employees of Dow Chemical and ExxonMobil, and two government employees. The commenter alleges that, in light of TERA's relationship with ExxonMobil, TERA's conclusions should be viewed with caution.

<u>CPSC Response 16</u>: We consider TERA to be an independent organization that focuses on advancing the science of toxicology and risk assessment. We do not agree that work by TERA or individual TERA staff in scientific projects, workshops, or publications concerning industrial chemicals or products or that include chemical firms, industry employees, or trade organizations necessarily indicates unreliable performance or improper influence in CPSC contract work.

As standard procedure, CPSC reviews potential conflicts of interest before awarding a contract or task order. We did not identify any conflicts for TERA related to the investigation of the production and use of phthalates or the production of the specified plastics. We do not agree that the membership in ARA is evidence of a potential conflict of interest. Rather, we consider ARA to be a transparent, multi-stakeholder scientific collaboration to develop risk assessment information to advance public health activities. Furthermore, the commenter does not specify any projects by the ARA that suggest that

¹ Prohibition of Children's Toys and Child Care Articles Containing Specified Phthalates: Determinations Regarding Certain Plastics, 8-26-17.

University of Cincinnati, College of Medicine

the contracted TERA work is affected by potential conflicts of interest.

Alachlor and Acetochlor 2

<u>Claim</u>: Dourson sought to undermine drinking water standards for the breakdown products of alachlor and acetochlor, two herbicides manufactured by Dow and Monsanto.

Reality:

- 1. TERA was approached by DowAgro Sciences and Monsanto to develop Reference Doses (RfDs) for degradates of these pesticides.
- 2. Michael Dourson talked with senior US EPA leaders to determine their interest.
- 3. EPA stated that they had developed RfDs for the parent chemicals and did not consider the degradates to be more toxic.
- 4. Michael Dourson suggested that DowAgro Sciences and Monsanto petition the Alliance for Risk Assessment (ARA) for their review.
- 5. The ARA Steering Committee endorsed a collaborative approach.
- TERA formed a team of risk assessment scientists from 3 states and the EPA to develop these RfDs. COIs statements were developed and reviewed at the meeting.
- 7. The meeting was open to the public.
- 8. The results were described in a report available to the public and in a publication.

1-Bromopropane³

<u>Claim</u>: TERA proposed a weaker standard for 1-bromopropane, a solvent used in degreasers, aerosol solvents, spray adhesives and dry cleaning.

Reality:

- 1. In 2004, occupation limits for 1-bromopropane differed by 16-fold.
- 2. TERA critically evaluated the underlying information and recommended an OEL of 20 ppm based on effects in newborns.
- 3. TERA's value was lower (i.e., safer) than EPA's.
- 4. An NTP study was conducted after the TERA assessment showing cancer findings.
- 5. New evaluations based on the cancer suggested lower limits.

- http://www.tera.org/ART/Degradates/index.html;
- Gadagbui, B; Maier, M; Dourson, M; Parker, A; Willis, A; Christopher, JP; Hicks, L; Ramasany, S; Roberts, SM. 2010. Derived Reference Doses (RfDs) for the Environmental Degradates of the Herbicides Alachlor and Acetochlor: Results of an Independent Expert Panel Deliberation. Regulatory Toxicology and Pharmacology 57:220-234.

² Source:

³ http://www.tera.org/Publications/TERA%20Analysis%20of%20OELs%20for%201-Bromopropane.pdf.

6. If confirmed, I will work with other federal agencies to develop a scientifically defensible position on this chemical under the LCSA.

Chlorpyrifos 4

<u>Claim</u>: Michael Dourson argued that chlorpyrifos was safe, despite three major studies showing that mothers and children who consume it are more at risk of giving birth to kids with ADHD and other neurological problems.

Reality:

- 1. TERA was funded by DowAgro Sciences to review the Reference Dose (RfD) developed by the EPA and others; results were published in 2005 and 2006.
- 2. The science for chlorpyrifos has progressed since the time of these publications.
- 3. One epidemiology study shows associations of neurological effects at exposures lower than the current RfD; other studies do not show this association.
- 4. Based on how chlorpyrifos works this association is not expected.
- 5. The raw data from this epidemiology study are not available for review.
- 6. If confirmed, I will work with investigators of this study to obtain these raw data, and will work with epidemiologists within EPA and other organizations to incorporate new information so that public health is protected.

Diacetyl 5

<u>Claim</u>: TERA sought to weaken standards for diacetyl, a chemical added to food and other products for flavor and aroma.

Reality:

- 1. At the time of TERA's work no standards existed for worker protection.
- 2. TERA's standard published in 2010 (i.e., range from 70 to 200 ppb) was based on the best science at the time, through careful consideration of toxicology, epidemiology, and background exposures.
- 3. Subsequent analyses published by various organizations include standards of 5 to 20 ppb based on different emphasis on toxicology and epidemiology data.
- 4. TERA is continuing its ongoing relationship with NIOSH since 2010 through an Interagency Personnel Agreement Fellowship.

• Z

- Zhao, Q., B. Gadagbui and M. Dourson. 2005. Lower birth weight as a critical effect of Chlorpyrifos: A comparison of human and animal data. Reg. Toxicol. Pharmacol. 42:55-63.
- Zhao, Q., M. Dourson and B. Gadagbui. 2006. A Review of the Reference Dose (RfD) for Chlorpyrifos. Reg. Toxicol. Pharmacol. 44:111-124.

⁴ Source:

⁵ Maier, AM; Kohrman-Vincent, M; Parker, A; Haber, LT. (2010) <u>Evaluation of concentration-response options for diacetyl in support of occupational risk assessment</u>. Reg. Toxicol. and Pharmacol. 58(2): 285-296.

5. This ongoing close relationship with TERA-NIOSH suggests that it finds TERA's work scientifically credible.

1,4-Dioxane 6

<u>Claim</u>: TERA sought to dramatically weaken the safety standard for 1,4-dioxane, an industrial chemical used in chemical processing.

Reality:

- 1. Dioxane occurs naturally in foods (up to 15 ppb in dairy products.
- 2. Dioxane causes cancer at high doses, but EPA's IRIS peer review panel thought that a nonlinear assessment might be appropriate.
- 3. Kentucky petitioned the Alliance for Risk Assessment to work collaboratively; 4 other states joined a request to the government of Japan, US NTP had previously helped.
- 4. Two publications resulted and support the EPA IRIS panel's nonlinear suggestion.
- 5. All of this information has been publicly available.
- 6. Health Canada is using TERA's collaborative work in their evaluation of dioxane.
- 7. If confirmed, I will work with other EPA offices to incorporate new information so that public health is protected.

Flame Retardants 7

<u>Claim</u>: Dourson served on Science Advisory Council of the North American Flame Retardant Alliance and co-wrote an article about the flame retardant chemical tetrabromobisphenol A, or TBBPA, casting doubt on whether the flame retardant has reproductive, neurological or development effects.

- Nishimura et al., 2004. Study of 1,4-dioxane intake in the total diet using the market-basket method. Journal of Health Science 50:101-107.
- Dourson, M; Reichard, J; Nance, P; Burleigh-Flayer, H; Parker, A; Vincent, M; McConnell, EE; (2014). Mode of Action Analysis for Liver Tumors from Oral 1,4-Dioxane Exposures and Evidence-Based Dose Response Assessment. Reg. Toxicol. Pharmacol. Volume 68, Issue 3, April 2014, Pages 387–401
- Michael L. Dourson, Jeri Higginbotham, Jeff Crum, Heather Burleigh-Flayer, Patricia Nance, Norman D. Forsberg, Mark Lafranconi, John Reichard. 2017. Update: Mode of action (MOA) for liver tumors induced by oral exposure to 1,4-dioxane. Regulatory Toxicology and Pharmacology 88:45-55.
- Website is currently in transfer mode. For current version see: http://med.uc.edu/eh/centers/rsc/risk-resources/ara.

⁶ Source:

⁷ Cope, Rhian B., Sam Kacew, Michael Dourson. 2015. A reproductive, developmental and neurobehavioral study following oral exposure of tetrabromobisphenol A on Sprague-Dawley rats. Toxicology 329 (2015) 49–59.

Reality:

- 1. Flame retardants save lives and property in innumerable situations. Countless examples exist of damage to property and lives lost when such chemicals are not available.
- 2. The Science Advisory Council of the NAFRA recommended publishing toxicology studies on several flame retardant chemicals so that the information was more publicly available, since current studies had been submitted to EPA in a confidential manner.
- 3. The publication on TBBPA showed no human relevant effects even at the highest dose used. This information can be used along with other toxicology studies to determine EPA's Reference Dose (RfD) for this chemical, which will then allow its regulation.
- 4. Michael Dourson worked with NAFRA so that this study, and all of its raw data, could be sent to the US NIEHS for their deliberation on whether to conduct a replicate study, thus potentially saving the US government about a million dollars.

Kids Chemical Safety website 8

<u>Claim</u>: Michael Dourson's TERA was given money by industry to create a misleading website on chemical safety for children.

Reality:

- 1. Stories on this kids website are written by identified experts for parents in an easier to understand way, since government websites are data-dense and activist websites appear designed for fundraising.
- 2. Experts are from <u>Cincinnati Children's Drug & Poison Information Center</u>, <u>Harvard Superfund Research Program</u>, <u>NSF International</u>, and <u>Toxicology Excellence for Risk Assessment (TERA)</u>.
- 3. TERA received cash gifts from the Alliance for Risk Assessment (*ARA*), American Chemistry Council (ACC), Combined Federal Campaign (CFC) of the US Federal Government, and the public.
- 4. Another nonprofit organization is reviewing this website for adoption.

MCHM-West Virginia 9

<u>Claim</u>: Michael Dourson did not disclose a conflict of interest prior to chairing this panel meeting.

http://web.archive.org/web/20161031132803/http://kidschemicalsafety.org/health/about/

⁸ Source:

⁹ Report of Expert Panel Review of Screening Levels for Exposure to Chemicals from the January 2014 Elk River Spill. West Virginia Testing Assessment Project, May 5, 2014.

Reality:

- 1. As for all of TERA's peer review meetings a COI disclosure was done prior to the meeting and commented on by all panel members.
- 2. This disclosure was part of the panel report.
- 3. West Virginia requested a closed review meeting, so such disclosures were not publicly available until the time of the press release the day after the meeting.
- 4. The Dourson-lead panel recommended the level of MCHM (4-methyl-1-cyclohexanemethanol) to be 8-fold more protective.
- 5. All of this information has been publicly available.

Peer Review 10

<u>Claim</u>: Over 50% of TERA's peer reviews are for industry. TERA whitewashes industry risk assessment values and places them on websites with other government information.

Reality:

- 1. Over 50% of TERA public peer review meetings were for governments.
- 2. Over 99% of TERA letter peer reviews were for governments.
- 3. All members of TERA's peer review panels were vetted for COI and balance was maintained among scientific disciplines and sector representation.
- 4. The panels decide whether information is sufficiently credible to load on the website.
- 5. The EPA IG (2009) commented favorably on TERA's peer review process, including its COI disclosures, COI updates at the meeting, and its documentation of COI in panel reports.
- 6. TERA is the only group to document COI decisions in its reports out of 6 groups reviewed by the EPA IG, including EPA's IRIS and the National Academy of Sciences.

Perchlorate 11

<u>Claim</u>: Michael Dourson's TERA was supported and paid to bless a high level of the rocket fuel perchlorate found at numerous sites around the country.

Reality:		

- http://www.tera.org/Peer/MeetingReports/index.html
- U.S. Environmental Protection Agency. 2009. Office Of Inspector General. EPA Can Improve Its Process For Establishing Peer Review Panels. Report No. 09-P-0147. April 29.

¹⁰ Source:

¹¹ Source: Strawson, J., Q. Zhao and M. Dourson. 2004. <u>Reference dose for perchlorate based on thyroid hormone change in pregnant women as the critical effect.</u> Reg. Tox. Pharm. 39: 44-65.

- 1. In 1995, PSG hired TERA to develop a safe dose, after EPA rejected PPG's level.
- 2. TERA developed a safe dose that was 100-fold lower (more protective), and recommended peer review.
- 3. The peer review recommended additional studies, which probably cost over 10 million dollars.
- 4. Afterwards EPA and the DOD disagreed on the safe dose.
- 5. TERA independently made its safe dose 5-fold more protective and published it.
- 6. The NAS also developed a safe dose, which was 25 times higher than EPA's, 12-fold lower than DoD's, but within 3 fold of TERA's value.
- 7. If confirmed, I will work with other EPA offices to incorporate new information so that public health is protected.

Petcoke-Chicago 12

<u>Claim</u>: Michael Dourson's TERA was supported by Koch industries to bless a petcoke storage facility in Chicago.

Reality:

- 1. The citizens of Chicago can make any risk management decision they desire regarding exposures to chemicals from any industry in their city.
- 2. TERA lead a team of scientists to determine exposures to petcoke in appropriate neighborhoods in Chicago.
- 3. Modeled exposures were compared to EPA's PM₁₀ NAAQS.
- 4. The work was published, allowing citizens of Chicago to consider these results in their risk management decision.

PFOA-Dupont 13

<u>Claim</u>: Michael Dourson's TERA was hand picked by Dupont and paid to bless a high level of PFOA in water in West Virginia.

Reality:

- 1. In 2002, 4 governments and one industry recommended TERA as the independent and neutral party to assist in a PFOA evaluation. A West Virginia judge agreed.
- 2. TERA, unaware of this agreement, was hired by the State of West Virginia.
- 3. Dr. Deanne Statts of West Virginia DEP chaired a 10-member scientific panel.
- 4. Five panelists were government employees; 3 were from EPA.
- 5. The panel made a unanimous determination of a safe water level of 150 ppb.
- 6. All of this information has been publicly available.

¹² Dourson, Michael, Chinkin, Lyle, MacIntosh, D.L., Finn, Jennifer, Brown, Kathleen, Reid, Stephen, Martinez, Jeanelle. 2016. A Case Study of Potential Human Health Impacts from Petroleum Coke Transfer Facilities. Journal of the Air & Waste Management Association May. DOI: 10.1080/10962247.2016.1180328

¹³ Source: FINAL CATT REPORT WITH ATTACHMENTS, AUGUST 2002

University of Cincinnati, College of Medicine

- 7. The science of PFOA has progressed since 2002.
- 8. If confirmed, I will work with other EPA offices to incorporate new information so that public health is protected.

Regulatory Toxicology and Pharmacology Journal 14

<u>Claim</u>: Michael Dourson publishes extensively in this journal, a mouthpiece of industry.

Reality:

- 1. This journal is unique in that it publishes papers that integrate toxicology and pharmacology findings into risk assessment and regulatory positions.
- 2. Because of this, many scientists from around the world publish in it.
- 3. My two most cited papers were in this journal.
- 4. I wrote these two papers as a US EPA employee.

Tobacco 15

Claim: Michael Dourson is a shill for the tobacco industry.

Reality:

- 1. TERA's work in tobacco has been previously described in a 2015 hearing of the U.S. House Committee on Science, Space, and Technology.
- 2. The total tobacco money received by TERA in 21 years was ~\$12,635.
- 3. Approximately \$6,000 was for a study on distribution of environmental tobacco smoke (ETS)-related chemicals for nonsmoking workers.
- 4. Approximately \$6,000 of this was for seminars on EPA's chemical mixtures risk assessment guidelines.
- 5. \$550 was for the development of a benchmark dose (BMD) for an ETS constituent, since the industry did not know how to use this new EPA method.
- 6. \$85 was for coping papers on work related to EPA's IRIS nickel document.

Trichloroethylene (TCE) 16

Michael Dourson, Bernard Gadagbui, Rod Thompson, Edward Pfau, and John Lowe.
 2016. Managing Risks of Noncancer Health Effects at Hazardous Waste Sites: A Case Study Using the Reference Concentration (RfC) of Trichloroethylene (TCE). Regulatory Toxicology and Pharmacology 80:125-133.

http://web.archive.org/web/20161031132803/http://kidschemicalsafety.org/health/about/

¹⁴ https://scholar.google.com/citations?user=N3DABAQAAAJ&hl=en

¹⁵ Response to Questions from U.S. House Committee on Science, Space, and Technology on EPA's 2015 Ozone Standard: Concerns Over Science and Implementation, Thursday, November 5, 2015.

¹⁶ Source:

<u>Claim</u>: Michael Dourson's TERA was hand picked by ACC and paid to bless a high level of TCE at superfund sites around the country.

Reality:

- 1. The Alliance for Risk Assessment (*ARA*) was petitioned by the Alliance for Site Closures to review noncancer toxicity of TCE.
- 2. The Steering Committee of the ARA, composed primarily of government officials, asked the collaboration to focus instead on building range in risk values.
- 3. The collaboration team had 6 conference calls, including scientists from Australia, 3 webinars, one of which included over 400 folks, and 1 independent peer consultation.
- 4. The team gave 8 presentations, and wrote one publication.
- 5. The team is planning training sessions with US states.
- 6. All of this information has been publicly available.
- 7. If confirmed, I will work with other EPA offices to incorporate new information so that public health is protected.

To: Hanley, Mary[Hanley.Mary@epa.gov]

Cc: Beck, Nancy[beck.nancy@epa.gov]; Bertrand, Charlotte[Bertrand.Charlotte@epa.gov]

From: Dourson, Michael

Sent: Thur 11/30/2017 12:17:13 AM

Subject: RE: PCBs in Schools 1-pager by Nov 30

Mary

Thanks for this information. Did the OPPT team also address the risk question regarding the 50 ppm level?

Cheers!

Michael

From: Hanley, Mary

Sent: Wednesday, November 29, 2017 6:16 PM

To: Beck, Nancy <Beck.Nancy@epa.gov>; Bertrand, Charlotte <Bertrand.Charlotte@epa.gov>;

Dourson, Michael <dourson.michael@epa.gov>
Subject: Fwd: PCBs in Schools 1-pager by Nov 30

Nancy,

Attached please find the PCBs in schools write-up that you requested from OPPT.

M

Sent from my iPhone

Begin forwarded message:

From: "Scheifele, Hans" < Scheifele. Hans@epa.gov>

Date: November 29, 2017 at 2:13:06 PM EST
To: "Hanley, Mary" < Hanley. Mary@epa.gov>
Cc: "Pierce, Alison" < Pierce. Alison@epa.gov>
Subject: RE: PCBs in Schools 1-pager by Nov 30

Mary,

Attached is a one-pager on PCBs that NPCD developed. Per your note below, Nancy asked for this by tomorrow. Please share with her and let me know if any questions.

Thanks,

Hans

Hans Scheifele Special Assistant Office of Pollution Prevention and Toxics 1200 Pennsylvania Ave., NW Washington, DC 20460 Voice (202) 564-3122

From: Scheifele, Hans

Sent: Tuesday, November 21, 2017 11:03 AM
To: Hanley, Mary < Hanley. Mary@epa.gov>
Cc: Pierce, Alison < Pierce. Alison@epa.gov>
Subject: Re: PCBs in Schools 1-pager by Nov 30

Hi Mary,

I'm out this week but Erik Winchester of NPCD is working on a response and Chris will

Ex. 5 - Deliberative Process

Hans

Sent from my iPhone

On Nov 19, 2017, at 8:55 AM, Hanley, Mary < Hanley. Mary@epa.gov > wrote:

Hi Hans, good question. Before I check with NB can you circle back to your folks to see if they have even back of the envelop costs/benefits. I only ask because this issue has been considered over the years. Thanks.

Cheers

Mary

From: Scheifele, Hans

Sent: Friday, November 17, 2017 1:22 PM

To: Hanley, Mary **Cc:** Pierce, Alison

Subject: RE: PCBs in Schools 1-pager by Nov 30

Mary,

For the cost benefit analysis, what is being asked for? Since a timeframe is being requested for a proposed rule this would include the development of an economic analysis. I am not aware that such an analysis is already available for this future proposal.

Can you please clarify what is needed?

Thanks, Hans

Hans Scheifele Special Assistant Office of Pollution Prevention and Toxics 1200 Pennsylvania Ave., NW Washington, DC 20460 Voice (202) 564-3122

----Original Message-----From: Hanley, Mary

Sent: Friday, November 17, 2017 12:16 PM
To: Scheifele, Hans < Scheifele. Hans@epa.gov >
Cc: Pierce, Alison < Pierce. Alison@epa.gov >
Subject: PCBs in Schools 1-pager by Nov 30

Hans

By Nov 30 th would OPPT draft a 1-pager (no more than 1 page per NB) on PCBs in schools. Please include the following:

>>> - info on concerns in PCBs in schools (NY, CA)

>>> - what we are doing now in states and regions -history/timeline for

>>> rulemaking -costs/benefits of the proposed rule.

>>> - OPPT workload effort to get a proposed and final rule out.

>>> Thanks M To: Beck, Nancy[Beck.Nancy@epa.gov]

From: Dourson, Michael

Sent: Tue 10/31/2017 10:53:22 AM

Subject: Re: GenX/PFOA/PFOS 1-2 Pager Request from the OCSPP IO

Another good idea... yes, I will bring this up with her.

Sent from my iPhone

On Oct 30, 2017, at 7:19 PM, Beck, Nancy < Beck.Nancy@epa.gov > wrote:

FYI on the fact sheet—see below.

Since ORD has the lead for the overarching effort, perhaps asking Jennifer OZ to set up the briefing for you may be the best path forward? Then ORD can also get you up to speed on their entire plan..

Are you chatting with her this week?

Nancy

Nancy B. Beck, Ph.D., DABT

Deputy Assistant Administrator, OCSPP

P: 202-564-1273

M: Ex. 6 - Personal Privacy

beck.nancy@epa.gov

From: Strauss, Linda

Sent: Monday, October 30, 2017 7:07 PM

To: Bertrand, Charlotte < Bertrand. Charlotte@epa.gov >

Cc: Beck, Nancy <<u>Beck.Nancy@epa.gov</u>>; Wise, Louise <<u>Wise.Louise@epa.gov</u>>; Hanley, Mary <<u>Hanley.Mary@epa.gov</u>>; Dunton, Cheryl <<u>Dunton.Cheryl@epa.gov</u>>

Subject: Re: GenX/PFOA/PFOS 1-2 Pager Request from the OCSPP IO

They were due Oct 26 but I know that some other offices have not gotten them in. Thanks for any edits. I think Nancy said she had thoughts too.

Ex. 5 - Deliberative Process

Thanks, Charlotte.

Sent from my iPhone

On Oct 30, 2017, at 6:44 PM, Bertrand, Charlotte < Bertrand. Charlotte@epa.gov > wrote:

Hi-I do have a number of comments, thought I would specifically edit the document to reflect my recommendations. When do you need this back? Charlotte

From: Strauss, Linda

Sent: Monday, October 30, 2017 10:46 AM

To: Beck, Nancy < Beck. Nancy@epa.gov >; Wise, Louise < Wise. Louise@epa.gov >;

Bertrand, Charlotte < Bertrand. Charlotte @epa.gov >

Cc: Hanley, Mary < Hanley. Mary@epa.gov >; Dunton, Cheryl

<Dunton.Cheryl@epa.gov>

Subject: FW: GenX/PFOA/PFOS 1-2 Pager Request from the OCSPP IO

Nancy, I know you mentioned you would have some comments. Just don't want this to fall off the radar. I think getting back to the OW organizer this week would be OK. Thanks, Linda

From: Strauss, Linda

Sent: Thursday, October 26, 2017 2:40 PM

To: Beck, Nancy < beck.nancy@epa.gov >; Wise, Louise < Wise.Louise@epa.gov >;

Bertrand, Charlotte < Bertrand. Charlotte@epa.gov >

Cc: Dunton, Cheryl < <u>Dunton.Cheryl@epa.gov</u>>; Hanley, Mary

< Hanley . Mary @epa.gov>

Subject: FW: GenX/PFOA/PFOS 1-2 Pager Request from the OCSPP IO

I think this has a lot of good information in it. It differs a little from what was requested which was for each AA-ship to provide its inventory of activities and messages for internal use. This does that but looks more like an Agency-wide fact sheet.

OW coms director who is coordinating this says she'd rather just see what we have – since it's only for internal use. At a minimum, we should take out the last heading. I have not made any edits. Let me know what you think.

Thanks, Linda

From: Scheifele, Hans

Sent: Wednesday, October 25, 2017 5:26 PM **To:** Strauss, Linda <<u>Strauss.Linda@epa.gov</u>>

Cc: Pierce, Alison <Pierce.Alison@epa.gov>; Dunton, Cheryl

<<u>Dunton.Cheryl@epa.gov</u>>; Hanley, Mary <<u>Hanley.Mary@epa.gov</u>> **Subject:** GenX/PFOA/PFOS 1-2 Pager Request from the OCSPP IO

Linda,

CCD developed the attached paper on PFOA/PFOS and RAD and Jeff reviewed and concurred. Please take a look and let us know if any questions.

Thanks,

Hans

Hans Scheifele Special Assistant Office of Pollution Prevention and Toxics 1200 Pennsylvania Ave., NW Washington, DC 20460 Voice (202) 564-3122

From: Drinkard, Andrea

Sent: Thursday, October 19, 2017 12:06 PM

To: Grantham, Nancy < <u>Grantham.Nancy@epa.gov</u>>; Lincoln, Larry

<<u>Lincoln.Larry@epa.gov</u>>; Mattas-Curry, Lahne <<u>Mattas-Curry.Lahne@epa.gov</u>>;

Hubbard, Carolyn < Hubbard. Carolyn@epa.gov >; Bowles, Jack

<<u>Bowles.Jack@epa.gov</u>>; Hannon, Arnita <<u>Hannon.Arnita@epa.gov</u>>; Richardson, RobinH <<u>Richardson.RobinH@epa.gov</u>>; Senn, John <<u>Senn.John@epa.gov</u>>; Jones,

Enesta < <u>Jones. Enesta@epa.gov</u>>; Maguire, Megan < <u>Maguire. Megan@epa.gov</u>>;

Strauss, Linda < Strauss. Linda @epa.gov >; Wadlington, Christina

< <u>Wadlington.Christina@epa.gov</u>>; <u>Millett, John < Millett.John@epa.gov</u>>;

Nowotarski, Allison < nowotarski.allison@epa.gov >

Cc: Klasen, Matthew < Klasen.Matthew@epa.gov >; Wise, Allison

< Wise. Allison@epa.gov>

Subject: GenX/PFOA/PFOS Comms Weekly Meeting Follow Up

Hi all—

In follow up to our call yesterday, I wanted to provide some information on what you might include in your one/two pagers on past/current/upcoming actions in your office. Here are some suggested section headers, but please feel free to use whatever format works best for your office as long as you are answering the general "what are we doing" question:

Overview/Background – Historically, what work has been done on PFAS in x office?

Current Action – What work is happening now?

What's Next – What work does x office anticipate need to complete to learn more about PFAS, etc.?

Current Messages – What would you tell someone today, if they asked "Is my water safe to drink? Is the air clean to breathe? Are these chemicals harmful to my health?"

Please provide me with your one/two pagers by COB on October 26th.

I'm working on a shared folder/sharepoint site and will update you when I have something concrete on that front. And, lastly, thanks to the subset of you who are looking at the blurb for Nancy. I will share the updated version with everyone on this email.

-Andrea-

Andrea Drinkard

Communications Director

EPA Office of Water

Desk: 202.564.1601

Cell: Ex. 6 - Personal Privacy

To: Keigwin, Richard[Keigwin.Richard@epa.gov]

From: Dourson, Michael

Sent: Tue 12/5/2017 12:25:16 PM Subject: RE: SRA Final Slides

Rick

Very nice.

Michael

From: Keigwin, Richard

Sent: Tuesday, December 5, 2017 6:57 AM

To: Bertrand, Charlotte <Bertrand.Charlotte@epa.gov>; Beck, Nancy <Beck.Nancy@epa.gov>; Wise, Louise <Wise.Louise@epa.gov>; Dourson, Michael <dourson.michael@epa.gov> **Cc:** Keller, Kaitlin <keller.kaitlin@epa.gov>; Dinkins, Darlene <Dinkins.Darlene@epa.gov>;

Vogel, Dana <Vogel.Dana@epa.gov>; Miller, David <Miller.DavidJ@epa.gov>

Subject: FW: SRA Final Slides

Next week, Steve Nako from OPP's Health Effects Division will be giving a presentation at the Society of Risk Analysis. His presentation will focus on trends in toxicity-adjusted dietary exposure to organophosphates and n-methyl carbamates. The discussion focuses on a measure for EPA's Report on the Environment. The measure depicts the impacts of pesticide regulatory actions on pesticide residues in food and the resulting dietary exposures since the passage of the Food Quality Protection Act of 1996. The downward trend is the result, directly and indirectly, of use restrictions that have been imposed on these chemicals as well as the registration of new alternative pest control technologies and strategies.

Let us know if you have any comments. SRA has asked to receive the PowerPoint in advance of the presentation.

From: Sent: Subject:	Dourson, Michael Thur 12/14/2017 7:47:47 PM meet earlier			
Nancy and Charlotte				
Can we r	neet earlier than 5 pm today? I have a 6 pm dinner at a metro stop a wee bit away.			
Cheers!				
Michael.				
L. Do	urson, PhD., DABT, FATS, FSRA			
Senior A	dvisor to the Administrator			
U.S. Environmental Protection Agency				
dourson.	michael@epa.gov			
202-564-	2463			
www.epa	a.gov			

Beck, Nancy[beck.nancy@epa.gov]; Bertrand, Charlotte[Bertrand.Charlotte@epa.gov]

To:

From: Sent: Subject:	Jackson, Ryanjjackson.ryan@epa.govj Dourson, Michael Thur 10/26/2017 6:00:31 PM RE: RE:
·	some text that might be helpful. Of course, please feel free to improve it.
Michael	
	x. 6 - Personal Privacy

Ex. 6 - Personal Privacy

From: Jackson, Ryan

Sent: Thursday, October 26, 2017 1:22 PM

To: Dourson, Michael dourson.michael@epa.gov

Subject: RE: RE:

Yep.

From: Dourson, Michael

Sent: Thursday, October 26, 2017 1:20 PM **To:** Jackson, Ryan jackson.ryan@epa.gov

Subject: Re: RE:

Sorry, would now be ok?

Sent from my iPhone On Oct 26, 2017, at 12:24 PM, Jackson, Ryan < jackson.ryan@epa.gov> wrote: I walked over, but come around when you get a moment. From: Dourson, Michael Sent: Thursday, October 26, 2017 11:41 AM To: Jackson, Ryan < jackson.ryan@epa.gov> Cc: Beck, Nancy < Beck. Nancy@epa.gov >; Lyons, Troy < lyons.troy@epa.gov > Subject: Re: Ryan Sure, how can I help? Michael Sent from my iPad On Oct 26, 2017, at 10:58 AM, Jackson, Ryan < <u>jackson.ryan@epa.gov</u>> wrote: I have one more request on this. Ryan Jackson Chief of Staff

U.S. Environmental Protection Agency

Ex. 6 - Personal Privacy

ED 001803B 00003025-3

To: Greenwalt, Sarah[greenwalt.sarah@epa.gov]; Washington,

Valerie[Washington.Valerie@epa.gov]

From: Dourson, Michael

Sent: Mon 11/27/2017 3:05:30 PM

Subject: RE: Running Late

Sarah and Valerie

I am on annual leave today, but look forward to seeing both of you tomorrow.

Cheers!

Michael

----Original Message----From: Greenwalt, Sarah

Sent: Monday, November 27, 2017 8:45 AM

To: Washington, Valerie < Washington. Valerie@epa.gov> Cc: Dourson, Michael < dourson.michael@epa.gov>

Subject: Re: Running Late

Hey Valerie, I'm on travel today but look forward to seeing you tomorrow.

Sent from my iPhone

> On Nov 27, 2017, at 7:22 AM, Washington, Valerie < Washington. Valerie@epa.gov> wrote:

>

- > Good Morning,
- > See you soon.

>

> Sent from my iPhone

To: Jackson, Ryan[jackson.ryan@epa.gov]

From: Dourson, Michael

Sent: Sat 10/21/2017 8:39:48 PM

Subject: Ohio Farmer Visits

Ryan

My November schedule has me at HQ on 1-3, 6-9, 13-17, and in Ohio for the week of Thanksgiving and the following Monday (1st day of gun deer season). I am then in HQ for 27-Dec 1. Please let me know if this schedule needs to be changed.

I also talked with Justina Fugh regarding whether it was within EPA ethics if I met with farmers and other interested folks in Ohio while I am there the week of Thanksgiving, especially if I am paying my own travel. She did not see a problem but also suggested talking with other folks to see if such meetings would be helpful in general.

So could such meetings be helpful to you and Administrator Pruitt? I can also bounce this off of Nancy Beck.

Cheers!

Michael

Sent from my iPad

To: Jackson, Ryan[jackson.ryan@epa.gov]; Bowman, Liz[Bowman.Liz@epa.gov]

Cc: Lyons, Troy[lyons.troy@epa.gov]

From: Dourson, Michael

Sent: Wed 11/1/2017 5:33:03 PM

Subject: RE: EPA SAB

Ryan

Thanks....

Michael

From: Jackson, Ryan

Sent: Wednesday, November 1, 2017 1:29 PM

To: Dourson, Michael dourson, Michael dourson.michael@epa.gov>

Cc: Lyons, Troy < lyons.troy@epa.gov>

Subject: RE: EPA SAB

Ex. 5 - Deliberative Process

From: Dourson, Michael

Sent: Wednesday, November 1, 2017 1:23 PM **To:** Bowman, Liz < <u>Bowman, Liz@epa.gov</u>>

Cc: Jackson, Ryan <<u>jackson.ryan@epa.gov</u>>; Lyons, Troy <<u>lyons.troy@epa.gov</u>>

Subject: FW: EPA SAB

Liz

Ex. 5 - Deliberative Process

Cheers!
Michael
From: Niina Heikkinen [mailto:nheikkinen@eenews.net] Sent: Wednesday, November 1, 2017 12:49 PM To: Dourson, Michael <dourson.michael@epa.gov> Subject: EPA SAB</dourson.michael@epa.gov>
Hi,
I'm curious what your thoughts are on the changes at EPA's science advisory board, as a former member. I understand you are not in a position to publicly comment but happy to speak on background or off the record.
Thanks,
Niina Heikkinen
E&E News reporter
202-737-3715 (w)
413-687-1789 (c)
@nhheikkinen
Skype: niina.h.heikkinen
E&E NEWS

122 C Street NW 7th Floor Washington, DC 20001

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Energywire, Climatewire, Greenwire, E&E Daily, E&E News PM

To: Beck, Nancy[Beck.Nancy@epa.gov]

From: Dourson, Michael

Sent: Tue 10/31/2017 10:51:52 AM

Subject: Re: watercress

Leaves for salad; everything for stirfry...

Sent from my iPhone

On Oct 30, 2017, at 7:21 PM, Beck, Nancy < Beck.Nancy@epa.gov > wrote:

Hey- for the salad, just the leaves or do people also eat the stems? thanks for sharing!!

Nancy B. Beck, Ph.D., DABT

Deputy Assistant Administrator

Office of Chemical Safety and Pollution Prevention

P: 202-564-1273

M: Ex. 6 - Personal Privacy

beck.nancy@epa.gov

To: King, Marquea[King.Marquea@epa.gov]

From: Dourson, Michael

Sent: Thur 12/14/2017 6:38:14 PM

Subject: RE: General

Marquee

No worries. The conversation was well worth the wee bit of wait...

Michael

From: King, Marquea

Sent: Thursday, December 14, 2017 12:54 PM **To:** Dourson, Michael dourson.michael@epa.gov

Subject: General

Hey there,

Thanks again, great seeing you! Sorry about being tardy.

MDKing

Marquea D. King, Ph.D.Mailing Address:P: 202-564-3626William Jefferson ClintonF: 202-564-8450

Designated Federal Official/

King Marquea@epa.gov

Mail Code 7202M (4126-J)
Toxicologist

1200 Pennsylvania Ave, NW

US Environmental Protection Agency

Washington, DC 20460

OCSPP/OSCP/Scientific Advisory Panel

To: Bolen, Derrick[bolen.derrick@epa.gov]

From: Dourson, Michael

Sent: Wed 11/8/2017 4:47:49 PM

Subject: Re: Thanksgiving

Derrick

I plan to work from my farm house on Monday, Tuesday and wednesday of thanksgiving week, and then take the Friday off.

Cheers!

Michael

Sent from my iPad

> On Nov 8, 2017, at 8:46 AM, Bolen, Derrick <bolen.derrick@epa.gov> wrote:

`

> All-

>

> If everyone could please send me what days they will be off for thanksgiving that would be greatly appreciated.

>

- > Thank you,
- > Derrick Bolen

To: Beck, Nancy[Beck.Nancy@epa.gov]

From: Dourson, Michael

Sent: Sat 10/21/2017 8:23:33 PM

Subject: Fwd: "A1 Table: some of these sources should not be trusted. For example..."

Nancy

Sorry, you missed the first of my several emails because I got your email backwards. Maybe this introductory email will clarify things.

Mike

Sent from my iPad

Begin forwarded message:

From: dourson.michael@epa.gov

Date: October 21, 2017 at 10:50:32 AM EDT

To: nancy.beck@epa.gov

Subject: "A1 Table: some of these sources should not be trusted. For example..."

Nancy

Here are some notes from my reading of the pre-prioritization text. In general, I thought

Ex. 5 - Deliberative Process

Attached are some notes taken on this text. Obviously my comments are helpful only as much as I understand the overall process. Admittedly I am still learning this. Thus, perhaps as expected, my comments are limited to more toxicological principles.

I will be sending other notes as well. Still learning how to use this iPad...

Cheers!

Mike

Open my shared note:

A1 Table: some of these sources should not be trusted. For example...

Notes

Sent from my iPad

To: Dominguez, Alexander[dominguez.alexander@epa.gov]

From: Dourson, Michael

Sent: Wed 11/1/2017 5:30:07 PM

Subject: RE: Charleston Hearing - Panel Participants

Dear Alex

I would be happy to throw my hat into the ring if you need a board certified toxicologist to do some of the listening. However, I am not a climate scientist.

Cheers!

Michael...

... L. Dourson, PhD., DABT, FATS, FSRA

Senior Advisor to the Administrator

U.S. Environmental Protection Agency

dourson.michael@epa.gov

202-564-2463

www.epa.gov

From: Dominguez, Alexander

Sent: Tuesday, October 31, 2017 2:43 PM

To: Dominguez, Alexander dominguez.alexander@epa.gov; Ford, Hayley

<ford.hayley@epa.gov>; Greenwalt, Sarah <greenwalt.sarah@epa.gov>; Frye, Tony (Robert)

<frye.robert@epa.gov>; Falvo, Nicholas <falvo.nicholas@epa.gov>; Hupp, Millan <hupp.millan@epa.gov>; Lovell, Will (William) <lovell.william@epa.gov>; Rodrick, Christian <rodrick.christian@epa.gov>; Shimmin, Kaitlyn <shimmin.kaitlyn@epa.gov>; Feeley, Drew (Robert) < Feeley. Drew@epa.gov>; Lyons, Troy < lyons.troy@epa.gov>; Forsgren, Lee <Forsgren.Lee@epa.gov>; Palich, Christian <palich.christian@epa.gov>; Yamada, Richard (Yujiro) <yamada.richard@epa.gov>; Ringel, Aaron <ringel.aaron@epa.gov>; Gordon, Stephen <gordon.stephen@epa.gov>; Abboud, Michael <abboud.michael@epa.gov>; Wagner, Kenneth <wagner.kenneth@epa.gov>; Kelly, Albert <kelly.albert@epa.gov>; Sands, Jeffrey <sands.jeffrey@epa.gov>; White, Elizabeth <white.elizabeth@epa.gov>; Cory, Preston (Katherine) <Cory.Preston@epa.gov>; Bowman, Liz <Bowman.Liz@epa.gov>; Letendre, Daisy <letendre.daisy@epa.gov>; Dravis, Samantha <dravis.samantha@epa.gov>; Ferguson, Lincoln <ferguson.lincoln@epa.gov>; Bolen, Brittany <bolen.brittany@epa.gov>; Harlow, David <harlow.david@epa.gov>; McMurray, Forrest < mcmurray.forrest@epa.gov>; Traylor, Patrick <traylor.patrick@epa.gov>; Greaves, Holly <greaves.holly@epa.gov>; Dominguez, Alexander <dominguez.alexander@epa.gov>; Beck, Nancy <Beck.Nancy@epa.gov>; Wilcox, Jahan <wilcox.jahan@epa.gov>; Baptist, Erik <baptist.erik@epa.gov>; Chmielewski, Kevin <chmielewski.kevin@epa.gov>; Hewitt, James <hewitt.james@epa.gov>; Schwab, Justin <Schwab.Justin@epa.gov>; Darwin, Henry <darwin.henry@epa.gov>; Bennett, Tate <Bennett.Tate@epa.gov>; Konkus, John <konkus.john@epa.gov>; Dourson, Michael <dourson.michael@epa.gov>; Bolen, Derrick <bolen.derrick@epa.gov>; Fotouhi, David <Fotouhi.David@epa.gov>; Bodine, Susan <bodine.susan@epa.gov>; Jackson, Ryan <jackson.ryan@epa.gov>; Munoz, Charles <munoz.charles@epa.gov>; Darwin, Veronica <darwin.veronica@epa.gov>; Brown, Byron
 brown.byron@epa.gov>

Cc: Gunasekara, Mandy < Gunasekara. Mandy @epa.gov>

Subject: Charleston Hearing - Panel Participants

All –

As you are aware, November 28 and 29 EPA will hold a public hearing in Charleston, WV, on the proposed repeal of the Clean Power Plan. We are looking for volunteers to attend to help coordinate and serve as panel participants. Panel participation is primarily a listening position whereby we hear testimony from interested stakeholders. We would like to assemble a list fairly quickly in order to let the travel coordinators know and so everyone can plan accordingly. If you are interested and available please confirm with me by 12:00 PM this Friday.

Alex Dominguez

Policy Analyst to the Deputy Assistant Administrator

Office of Air and Radiation

U.S. Environmental Protection Agency

To: Beck, Nancy[Beck.Nancy@epa.gov]

From: Dourson, Michael

Sent: Tue 10/31/2017 10:46:11 AM

Subject: Re: OPP Weekly Report for Week Ending Friday, October 27th

Wow! Impressive...

Sent from my iPhone

On Oct 30, 2017, at 10:10 PM, Beck, Nancy < Beck.Nancy@epa.gov > wrote:

FYI

Nancy B. Beck, Ph.D., DABT
Deputy Assistant Administrator, OCSPP
P: 202-564-1273
M: Ex. 6 - Personal Privacy

From: Keller, Kaitlin

Beck.Nancy@epa.gov

Sent: Monday, October 30, 2017 6:11 PM

To: Beck, Nancy < <u>Beck.Nancy@epa.gov</u>>; Bertrand, Charlotte

<<u>Bertrand.Charlotte@epa.gov</u>>; Wise, Louise <<u>Wise.Louise@epa.gov</u>> **Subject:** FW: OPP Weekly Report for Week Ending Friday, October 27th

FYI. OPP Weekly Report attached.

From: Jewell, Shannon

Sent: Monday, October 30, 2017 5:09 PM **To:** OPP ALL < OPP ALL @epa.gov >

Subject: OPP Weekly Report for Week Ending Friday, October 27th

Hello All,

Please access the report by clicking the link and signing into SharePoint here: <u>OPP Weekly Report for Week Ending 10/27/17.</u>
Please remember that the OPP Weekly Report is for internal distribution only.
Thank you,
Shannon
Shannon Jewell · (703) 347-0109 · jewell.shannon@epa.gov
EPA Office of Pesticide Programs Immediate Office
<opp 10.27.17.pdf="" report="" weekly=""></opp>

To: Munoz, Charles[munoz.charles@epa.gov]

Cc: Washington, Valerie[Washington.Valerie@epa.gov]; Jenkins, Donna[Jenkins.Donna@epa.gov]

From: Dourson, Michael

Sent: Thur 1/4/2018 3:13:42 PM

Subject: Shipping

Charles

Ex. 6 - Personal Privacy

The addresses are all on the boxes, but just in case, they go to:

Michael L. Dourson

Ex. 6 - Personal Privacy

Thanks!

Michael

Ex. 6 - Personal Privacy (cell) (home)

From: Munoz, Charles

Sent: Tuesday, January 2, 2018 5:54 PM

Subject: Re: Talk
Yep. I'll be in no later than 8 so come by anytime between that and 10:15 if possible. I'm free after 10:45 till about 1 as well.
Charles Munoz
White House Liaison
On Jan 2, 2018, at 5:51 PM, Dourson, Michael < <u>dourson.michael@epa.gov</u> > wrote:
Charles
Can we talk tomorrow morning?
Cheers!
Michael
L. Dourson, PhD., DABT, FATS, FSRA
Senior Advisor to the Administrator
U.S. Environmental Protection Agency
dourson.michael@epa.gov
202-564-2463
www.epa.gov

To: Zarba, Christopher[Zarba.Christopher@epa.gov]

From: Dourson, Michael

Sent: Thur 12/14/2017 6:29:47 PM

Subject: Re: Misc.

Chris

Thanks for your kind words. The SAB is very important. Honeycutt is a good choice as lead.

Mike

Sent from my iPhone

On Dec 14, 2017, at 11:49 AM, Zarba, Christopher < Zarba. Christopher@epa.gov > wrote:

I am sorry to see you go. I was looking forward to working with you.

Best of luck in future endeavors and please stay in touch...

Christopher S. Zarba

US EPA Science Advisory Board

zarba.christopher@epa.gov

O (202) 564-0760

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